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(54) Title: METHODS FOR PRODUCING MEMBERS OF SPECIFIC BINDING PAIRS

(57) Abstract

A member of a specific binding pair (sbp) is identified by expressing DNA encoding a genetically diverse population of such sbp members in recombinant host cells in which the sbp members are displayed in functional form at the surface of a secreted recombinant genetic display package (rgdp) containing DNA encoding the sbp member or a polypeptide component thereof, by virtue of the sbp member or a polypeptide component thereof being expressed as a fusion with a capsid component of the rgdp. The displayed shps may be selected by affinity with a complementary shp member, and the DNA recovered from selected rgdps for expression of the selected sbp members. Antibody sbp members may be thus obtained, with the different chains thereof expressed, one fused to the capsid component and the other in free form for association with the fusion partner polypeptide. A phagemid may be used as an expression vector, with said capsid fusion helping to package the phagemid DNA. Using this method libraries of DNA encoding respective chains of such multimeric sbp members may he combined, thereby obtaining a much greater genetic diversity in the sbp members than could easily be obtained by conventional methods.





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METHODS FOR PRODUCING MEMBERS OF SPECIFIC BINDING PAIRS

The present invention relates to methods for producing members of specific binding pairs. The present invention also relates to the biological binding molecules produced by these methods.

Owing to their high specificity for a given antigen, the advent of monoclonal antibodies (Kohler, G. and Milstein 10 C; 1975 Nature 256: 495) represented a significant technical break-through with important consequences both scientifically and commercially.

Monocional antibodies are traditionally made by

establishing an immortal mammalian cell line which is 15 derived from a single immunoglobulin producing cell secreting one form of a biologically functional antibody molecule with a particular specificity. Because the antibody-secreting mammalian cell line is immortal, the characteristics of the antibody are reproducible from batch 20 to batch. The key properties of monoclonal antibodies are

their specificity for a particular antigen and the

reproducibility with which they can be manufactured.

Structurally, the simplest antibody (IgG) comprises four polypeptide chains, two heavy (H) chains and two light 25 (L) chains inter-connected by disulphide bonds (see figure The light chains exist in two distinct forms called kappa (K) and lambda (λ) . Each chain has a constant region (C) and a variable region (V). Each chain is organized into

a series of domains. The light chains have two domains. corresponding to the C region and the other to the V region. 30 The heavy chains have four domains, one corresponding to the V region and three domains (1,2 and 3) in the C region. The antibody has two arms (each arm being a Fab region), each of

which has a VL and a VH region associated with each other. It is this pair of V regions (VL and VH) that differ from 35 one antibody to another (owing to amino acid sequence variations), and which together are responsible for recognising the antigen and providing an antigen binding site (ABS). In even more detail, each V region is made up 40

from three complementarity determining regions (CDR) separated by four framework regions (FR). The CDR's are the most variable part of the variable regions, and they perform the critical antigen binding function. The CDR regions are derived from many potential germ line sequences via a 45

complex process involving recombination, mutation and selection.

It has been shown that the function of binding entigens can be performed by fragments of a whole antibody. Example binding fragments are (i) the Fab fragment consisting of the 50 VL, VH, CL and CH1 domains: (ii) the Fd fragment consisting of the VH and CH1 domains; (iii) the Fv fragment consisting of the VL and VH domains of a single arm of an antibody, (iv) the dAb fragment (Ward, E.S. et al., Nature 341, 544-546 (1989) which consists of a VH domain; (v) isolated CDR

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regions; and (vi) F(ab')2 fragments, a bivalent fragment comprising two Fab fragments linked by a disulphide bridge

at the hinge region.

Although the two domains of the Fv fragment are coded for by separate genes, it has proved possible to make a synthetic linker that enables them to be made as a single protein chain (known as single chain Fv (scFv); Bird, R.E. et al., Science 242, 423-426 (1988) Huston, J.S. et al., Proc. Natl. Acad. Sci., USA 85, 5879-5883 (1988)) by recombinant methods. These scrv fragments were assembled from genes from monoclonals that had been previously isolated. In this application, the applicants describe a process to assemble scFv fragments from VH and VL domains that are not part of an antibody that has been previously

15 isolated.

Whilst monoclonal antibodies, their fragments and derivatives have been enormously advantageous, there are nevertheless a number of limitations associated with them.

Firstly, the therapeutic applications of monoclonal antibodies produced by human immortal cell lines holds great 20 promise for the treatment of a wide range of diseases (Clinical Applications of Monoclonal Antibodies. Edited by E. S. Lennox. British Medical Bulletin 1984. Publishers Churchill Livingstone). Unfortunately, immortal antibody-

producing human cell lines are very difficult to establish 25 and they give low yields of antibody (approximately 1 ug/ml). In contrast, equivalent rodent cell lines yield high amounts of antibody (approximately 100 µg/ml). However, the repeated administration of these foreign rodent proteins to humans can lead to harmful hypersensitivity reactions. In the main therefore, these 30

rodent-derived monoclonal antibodies have limited

therapeutic use.

Secondly, a key aspect in the isolation of monoclonal antibodies is how many different clones of antibody 35 producing cells with different specificities, can be practically established and sampled compared to how many theoretically need to be sampled in order to isolate a cell producing antibody with the desired specificity characteristics (Milstein, C., Royal Soc. Croonian Lecture, Proc. R. Soc. London B. 239; 1-16, (1990)). For example, the number of different specificities expressed at any one 40 time by lymphocytes of the murine immune system is thought to be approximately 10^7 and this is only a small proportion

of the potential repertoire of specificities. However, 45 during the isolation of a typical antibody producing cell with a desired specificity, the investigator is only able to sample 10^3 to 10^4 individual specificaties. The problem is worse in the human, where one has approximately 1012 lymphocyte specificities, with the limitation on simpling of $10^3 \ \rm or \ 10^4 \ \rm remaining$. 50

This problem has been alleviated to some extent in laboratory animals by the use of immunisation regimes. Thus, where one wants to produce monoclonal antibodies having a specificity against a particular epitope, an animal 55

is immunised with an immunogen expressing that epitope. The animal will then mount an immune response against the immunogen and there will be a proliferation of lymphocytes which have specificity against the epitope. Owing to this proliferation of lymphocytes with the desired specificity. it becomes easier to detect them in the sampling procedure. However, this approach is not successful in all cases, as a suitable immunogen may not be available. Furthermore, where one wants to produce human monoclonal antibodies (eg for therapeutic administration as previously discussed), such an 10 approach is not practically, or ethically, feasible.

In the last few years, these problems have in part, been addressed by the application of recombinant DNA methods to the isolation and production of e.g. antibodies and fragments of antibodies with antigen binding ability, in

bacteria such as E.coli. This simple substitution of immortalised cells with bacterial cells as the 'factory', considerably simplifies procedures for preparing large amounts of binding molecules. Furthermore, a recombinant production system allows scope 20 for producing tailor-made antibodies and fragments thereof. For example, it is possible to produce chimaeric molecules with new combinations of binding and effector functions, humanised antibodies (e.g. murine variable regions combined 25 with human constant domains or murine-antibody CDRs grafted onto a human FR) and novel antigen-binding molecules. Furthermore, the use of polymerase chain reaction (PCR) amplification (Saiki, R.K., et al., Science 239, 487-491

(1988)) to isolate antibody producing sequences from cells (e.g. hybridomas and B cells) has great potential for 30 speeding up the timescale under which specificities can be isolated. Amplified VH and VL genes are cloned directly into vectors for expression in bacteria or mammalian cells (Orlandi, R., et al., 1989, Proc. Natl. Acad. Sci., USA 86,

35 3833-3837; Ward, E.S., et al., 1989 supra: Larrick, J.W., et al., 1989, Biochem. Biophys. Res. Commun. 160, 1250-1255; Sastry, L. et al., 1989, Proc. Natl. Acad. Sci., USA., 86, 5728-5732). Soluble antibody fragments secreted from bacteria are then screened for binding activities.

40 However, like the production system based upon

immortalised cells, the recombinant production system still suffers from the selection problems previously discussed and therefore relies on animal immunization to increase the proportion of cells with desired specificity. Furthermore, some of these techniques can exacerbate the screening 45 For example, large separate H and L chain problems. libraries have been produced from immunized mice and combined together in a random combinatorial manner prior to screening (Huse, W.D. et al., 1989, Science 246, 1275-1281,

50 W090/14443; W090/14424 and W090/14430). Crucially however, the information held within each cell, namely the original pairing of one L chain with one H chain, is lost. This loses some, of the advantage gained by using immunization protocols in the animal. Currently, only libraries derived from single VH domains (dAbs; Ward, E.S., et al., 1989, 55

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supra.) do not suffer this drawback. However, because not all antibody VH domains are capable of binding antigen, more have to be screened. In addition, the problem of directly screening many different specificities in prokaryotes remains to be solved.

Thus, there is a need for a screening system which ameliorates or overcomes one or more of the above or other problems. The ideal system would allow the sampling of very large numbers of specificities (eg 10^6 and higher), rapid sorting at each cloning round, and rapid transfer of the genetic material coding for the binding molecule from one

stage of the production process, to the next stage.

The most attractive candidates for this type of screening, would be prokaryotic organisms (because they grow quickly, are relatively simple to manipulate and because large numbers of clones can be created) which express and display at their surface a functional binding domain eg. an In the UK patent GB antibody, receptor, enzyme etc. 21376318 methods for the co-expression in a single host cell of the variable H and L chain genes of immunoglobulins were However, the protein was expressed disclosed. intracellularly and was insoluble. Further, the protein required extensive processing to generate antibody fragments with binding activity and this generated material with only a fraction of the binding activity expected for antibody fragments at this concentration. It has already been shown that antibody fragments can be secreted through bacterial

membranes with the appropriate signal peptide (Skerra, A. and Pluckthun, A. 1988 Science 240 1038-1040; Better, M et al 1988, Science 240 1041-1043) with a consequent increase in the binding activity of antibody fragments. methods require screening of individual clones for binding activity in the same way as do mouse monoclonal antibodies.

It has not been shown however, how a functional binding domain eg an antibody, antibody fragment, receptor, enzyme 35 etc can be held on the bacterial surface in a configuration which allows sampling of say its antigen binding properties and selection for clones with desirable properties. large part, this is because the bacterial surface is a complex structure, and in the gram-negative organisms there 40 is an outer wall which further complicates the position.

Further, it has not been shown that eg an antibody domain will fold correctly when expressed as a fusion with a

surface protein of bacteria or bacteriophage.

Bacteriophage are attractive prokaryote related 45 organisms for this type of screening. In general, their surface is a relatively simple structure, they can be grown easily in large numbers, they are amenable to the practical handling involved in many potential mass screening programmes, and they carry genetic information for their own 50

synthesis within a small, simple package. The difficulty has been to practically solve the problem of how to use bacteriophages in this manner. A Genex Corporation patent application number WO88/06630 has proposed that the bacteriophage lambda would be a suitable vehicle for the 45

expression of antibody molecules, but they do not provide a teaching which enables the general idea to be carried out. For example W088/06630 does not demonstrate that any sequences: (a) have been expressed as a fusion with gene V; (b) have been expressed on the surface of lambda; and (c) have been expressed so that the protein retains biological activity. Furthermore there is no teaching on how to screen for suitable fusions. Also, since the lambda virions are assembled within the cell, the fusion protein would be 10 expressed intracellularly and would be predicted to be inactive. Bass et al., in December 1990 (after the earliest priority date for the present application) describe deleting part of gene III of the filamentous bacteriophage M13 and inserting the coding sequence for human growth hormone (hGH) 15 into the N-terminal site of the gene. The growth hormone displayed by M13 was shown to be functional. (Bass, S., et al. Proteins, Structure, Function and Genetics (1990) 8:

309-314). A functional copy of gene III was always present in addition, when this fusion was expressed. A Protein Engineering Corporation patent application W090/02809 proposes the insertion of the coding sequence for bovine pancreatic trypsin inhibitor (BPTI) into gene VIII of M13. However, the proposal was not shown to be operative. For example, there is no demonstration of the expression of BPTI

25 sequences as fusions with protein VIII and display on the surface of Mi3. Furthermore this document teaches that when a fusion is made with gene III, it is necessary to use a second synthetic copy of gene III, so that some unaltered gene III protein will be present. The embodiments of the present application do not do this. In embodiments where phagemid is rescued with Mi3KO7 gene III deletion phage,

there is no unaltered gene III present.

W090/02809 also teaches that phagemids that do not contain the full genome of M13 and require rescue by coinfection with helper phage are not suitable for these purposes because coinfection could lead to recombination.

In all embodiments where the present applicants have used phagemids, they have used a helper phage and the only sequences derived from filamentous bacteriophage in the phagemids are the origin of replication and gene III

sequences.

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W090/02809 also teaches that their process needed information such as nucleotide sequence of the starting molecule and its three-dimensioned structure. The use of a pre-existing repertoire of binding molecules to select for a binding member, such as is disclosed herein, for example using an immunoglobulin gene repertoire of animals, was not disclosed. Further, they do not discuss favouring variegation of their binding molecules in natural blocks of variation such as CDRs of immunoglobulins, in order to revolve generation of improved molecules and prevent unfavourable variations. W090/02809 also specifically excluded the application of their process to the production of scfv molecules.

In each of the above discussed patents (WO88/O6630 and

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W090/02809), the protein proposed for display is a single There is no disclosure of a method for polypeptide chain. the display of a dimeric molecule by expression of one monomer as a fusion with a capsid protein and the other protein in a free form.

Another disclosure published in May 1991 (after the earliest priority date for the present application) describes the insertion into gene VIII of M13, the coding sequences for one of the two chains of the Fab portion of an antibody with co-expression of the other from a plasmid. The two chains were demonstrated as being expressed as a

functional Fab fragment on the surface of the phage (Kang A.S. et al., (1991) Proc. Natl. Acad. Sci, USA, 88 p4363-4366). No disclosure was made of the site of insertion into gene VIII and the assay for pAb binding activity by ELISA used a reagent specific for antibody L chain rather than for phage. A further disclosure published in March 1991 (after the earliest priority date for the present application) describes the insertion of a fragment of the AIDS virus protein gag into the N-terminal portion of gene III of the bacteriophage fd. The expression of the gag protein fragment was detected by immunological methods, but it was not shown whether or not the protein was expressed in a

functional form (Tsunetsugu-Yokota Y et al. (1991) Gene 99 p261-265).

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The problem of how to use bacteriophages in this way is in fact a difficult one. The protein must be inserted into the phage in such a way that the integrity of the phage coat is not undermined, and the protein itself should be functional retaining its biological activity with respect to 30 antigen binding. Thus, where the protein of choice is an antibody, it should fold efficiently and correctly and be presented for antigen binding. Solving the problem for antibody molecules and fragments would also provide a general method for any biomolecule which is a member of a 35 specific binding pair e.g. receptor molecules and enzymes.

Surprisingly, the applicants have been able to construct a bacteriophage that expresses and displays at its surface a large biologically functional binding molecule (eg antibody fragments, and enzymes and receptors) and which remains intact and infectious. The applicants have called 40 the structure which comprises a virus particle and a binding molecule displayed at the viral surface a 'package'. Where the binding molecule is an antibody, an antibody derivative or fragment, or a domain that is homologous to an 45 immunoglobulin domain, the applicants call the package a

'phage antibody' (pAb). However, except where the context demands otherwise, where the term phage antibody is used generally, it should also be interpreted as referring to any package comprising a virus particle and a biologically functional binding molecule displayed at the viral surface.

pAbs have a range of applications in selecting antibody genes encoding antigen binding activities. For example, pAbs could be used for the cloning and rescue of hybridomas 55

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(Orlandi, R., et al (1989) PNAS 86 p3833-3837), and in the screening of large combinatorial libraries (such as found in Huse, W.D. et al., 1989, Science 246, 1275-1281). particular, rounds of selection using pabs may help in rescuing the higher affinity antibodies from the latter libraries. It may be preferable to screen small libraries derived from antigen-selected cells (Casali, P., et al., (1986) Science 234 p476-479) to rescue the original VH/VL pairs comprising the Fv region of an antibody. The use of pabs may also allow the construction of entirely synthetic antibodies. Furthermore, antibodies may be made which have some synthetic sequences e.g. CDRs, and some naturally derived sequences. For example, V-gene repertoires could be made in vitro by combining un-rearranged V genes, with D and Libraries of pabs could then be selected by J segments. binding to antigen, hypermutated in vitro in the antigenbinding loops or V domain framework regions, and subjected to further rounds of selection and mutagenesis. As previously discussed, separate H and L chain

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20 libraries lose the original pairing between the chains. It is difficult to make and screen a large enough library for a particularly advantageous combination of H and L chains.

For example, in a mouse there are approximately 10⁷ possible H chains and 10⁷ possible L chains. Therefore, there are 10¹⁴ possible combinations of H and L chains, and to test for anything like this number of combinations one would have to create and screen a library of about 10¹⁴ clones. This has not previously been a practical possibility.

30 The present invention provides a number of approaches which ameliorate this problem.

In a first approach, (a random combinatorial approach, see examples 20 and 21) as large a library as is practically possible is created which expresses as many of the 10¹⁴ potential combinations as possible. However, by virtue of the expression of the H and L chains on the surface of the phage, it is reasonably practicable to select the desired combination, from all the generated combinations by affinity techniques (see later for description of selection formats).

techniques (see later for description of selection formats).

In a second approach (called a dual combinatorial approach by the present applicants, see example 26), a large library is created from two smaller libraries for selection of the desired combination. This ameliorates the problems still further. The approach involves the creation of: (i) a first library of say 10° e.g. H chains which are displayed on a bacteriophage (as a fusion with the protein encoded by gene III) which is resistant to e.g. tetracycline: and (ii) a second library of say 10° e.g. L chains in which the coding sequences for these light chains are within a plasmid vector containing an origin of replication for a bacteriophage (a phagemid) which is resistant to e.g. ampicillin (i.e. a different antibiotic) and are expressed in the periplasmic space of a host bacterium. The first

library is then used to infect the bacteria containing the 55 second library to provide 10^{14} combinations of 8 and L

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chains on the surface of the resulting phage in the

bacterial supernatant.

The advantage of this approach is that two separate libraries of eg 10^7 are created in order to produce 10^{14} combinations. Creating a 10^7 library is a practical

possibility. The 10^{14} combinations are then subjected to selection (see later for description of selection formats) as disclosed by the present application. This selection will then produce a population of phages displaying a particular 10 combination of H and L chains having the desired Specificity. The phages selected however, will only contain DNA encoding one partner of the paired H and L chains (deriving from either the phage or phagemid). The sample eluate containing the population is then divided into two 15 portions. A first portion is grown on e.g. tetracycline plates to select those bacteriophage containing DNA encoding H chains which are involved in the desired antigen binding. A second portion is grown on e.g. ampicillin plates to select those bacteriophage containing phagemid DNA encoding 20 L chains which are involved in the desired antigen binding. A set of colonies from individually isolated clones e.g. from the tetracycline plates are then used to infect specific colonies e.g. from the ampicillin plates. results in bacteriophage expressing specific combinations of 25 H and L chains which can then be assayed for antigen

binding.

In a third approach (called a hierarchical dual combinational approach by the present applicants), an individual colony from either the H or L chain clone selected by growth on the antibiotic plates, is used to infect a complete library of clones encoding the other chain (H or L). Selection is as described above. This favours isolation of the most favourable combination.

In a fourth approach (called a hierarchrical approach by the present applicants, see examples 22 and 46) both chains are cloned into the same vector. However, one of the chains which is already known to have desirable properties is kept fixed. A library of the complementary chain is inserted into the same vector. Suitable partners for the fixed chain are selected following display on the surface of bacteriophage.

In a fifth approach (see example 48), to improve the chances of recovering original pairs, the complexity of the combinatorial libraries can be reduced by using small be populations of B-lymphocytes selected for binding to a desired antigen. The cells provide e.g. mRNA or DNA, for preparing libraries of antibody genes for display on phage. This technique can be used in combination with the above mentioned four approaches for selection of antibody specificities.

Phagemids have been mentioned above. The applicants have realised and demonstrated that in many cases phagemids will be preferred to phage for cloning antibodies because it is easier to use them to generate more comprehensive

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libraries of the immune repertoire. This is because the phagemid DNA is approximately 100 times more efficient than bacteriophage DNA in transforming bacteria (see example 19). Also, the use of phagemids gives the ability to vary the number of gene III binding moecule fusion proteins displayed on the surface of the bacteriophage (see example 17). For example, in a system comprising a bacterial cell containing a phagemid encoding a gene III fusion protein and infected with a helper phage, induction of expression of the gene III fusion protein to different extents, will determine the number of gene III fusion proteins present in the space defined between the inner and outer bacterial membranes following superinfection. This will determine the ratio of gene III fusion protein to native gene III protein displayed

15 by the assembled phage.

Expressing a single fusion protein per virion may aid selection of antibody specificities on the basis of affinity by avoiding the 'avidity' effect where a phage expressing two copies of a low affinity antibody would have the same apparent affinity as a phage expressing one copy of a higher affinity antibody. In some cases however, it will be important to display all the gene III molecules derived by superinfection of cells containing phagemids to have fusions (e.g. for selecting low affinity binding molecules or improving sensitivity on ELISA). One way to do this is to

25 improving sensitivity on ELISA). One way to do this is to superinfect with a bacteriophage which contains a defective gene III. The applicants have therefore developed and used a phage which is deleted in gene III. This is completely novel.

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The demonstration that a functional antigen-binding domain can be displayed on the surface of phage, has implications beyond the construction of novel antibodies. For example, if other protein domains can be displayed at the surface of a phage, phage vectors could be used to clone and select genes by the binding properties of the displayed protein. Furthermore, variants of proteins, including epitope libraries built into the surface of the protein, could be made and readily selected for binding activities. In effect, other protein architectures might serve as

40 "nouvelle" antibodies.

The technique provides the possibility of building antibodies from first principles, taking advantage of the structural framework on which the antigen binding loops fold. In general, these loops have a limited number of conformations which generate a variety of binding sites by alternative loop combinations and by diverse side chains. Recent successes in modelling antigen binding sites sugurs well for de novo design. In any case, a high resolution structure of the antigen is needed. However, the approach is attractive for making e.g. catalytic antibodies, particularly for small substrates. Here side chains or binding sites for prosthetic groups might be introduced, not only to bind selectively to the transition state of the substrate, out also to participate directly in bond making and breaking. The only question is whether the antibody

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architecture, specialised for binding, is the best starting point for building catalysts. Genuine enzyme architectures, such as the triose phosphate isomerase (TIM) barrel, might be more suitable. Like antibodies, TIM enzymes also have a framework structure (a barrel of β -strands and α -helices) and loops to bind substrate. Many enzymes with a diversity of catalytic properties are based on this architecture and the loops might be manipulated independently on the frameworks for design of new catalytic and binding The phage selection system as provided by the properties. present disclosure can be used to select for antigen binding activities and the CDR loops thus selected, used on either an antibody framework or a TIM barrel framework. Loops placed on a e.g. a TIM barrel framework could be further modified by mutagenesis and subjected to further selection. Thus, there is no need to select for high affinity binding activities in a single step. The strategy of the immune

more realistic and can be mimicked using this invention. One class of molecules that could be useful in this type of application are receptors. For example, a specific receptor could be displayed on the surface of the phage such that it would bind its ligand. The receptor could then be modified by, for example, in vitro mutagenesis and variants having higher binding affinity for the ligand selected. The selection may be carried out according to one or more of the formats described below with reference to figure 2 (which refers particularly to pAbs) in which the pAb antibody is

system, in which low affinity evolves to high affinity seems

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replaced with a phage receptor and the antigen with a ligand Alternatively, the phage-receptor could be used as the basis of a rapid screening system for the binding of ligands, altered ligands, or potential drug candidates. The advantages of this system namely of simple cloning, convenient expression, standard reagents and easy handling 35 makes the drug screening application particularly In the context of this discussion, receptor attractive. means a molecule that binds a specific, or group of specific, ligand(s). The natural receptor could be expressed on the surface of a population of cells, or it could be the extracellular domain of such a molecule (whether such a form exists naturally or not), or a soluble molecule performing a natural binding function in the plasma, or within a cell or organ.

45 Another possibility, is the display of an enzyme molecule or active site of an enzyme molecule on the surface of a phage (see examples 11,12,30,31,32 and 36). Once the phage enzyme is expressed, it can be selected by affinity chromatography, for instance on columns derivatized with transition state analogues. If an enzyme with " different or modified specificity is desired, it may be possible to mutate an enzyme displayed as a fusion on bacteriophage and then select on a column derivatised with an analogue selected to have a higher affinity for an enzyme with the

55 desired modified specificity.

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Although throughout this application, the applicants discuss the possibility of screening for higher affinity variants of pAbs, they recognise that in some applications, for example low affinity chromatography (Ohlson, S. et al Anal. Biochem. 169, p204-208 (1988)) it may be desirable to isolate lower affinity variants.

Examples 21 and 23 show that the present invention provides a way of producing antibodies with low affinities (as seen in the primary immune response or in 1.0 unimmunised animals). This is made possible by displaying multiple copies of the antibody on the phage surface in association with gene III protein. Thus, pabs allow genes for these antibodies to be isolated and if necessary, mutated to provide improved antibodies.

15 pAbs also allow the selection of antibodies for improved stability. It has been noted for many antibodies, that yield and stability are improved when the antibodies are expressed at 30°C rather than 37°C. If pAbs are displayed at 37°C, only those which are stable will be available for affinity selection. antibodies are to be used in vivo for therapeutic or diagnostic purposes, increased stability would extend the

half-life of antibodies in circulation. Although stability is important for all antibodies 25 and antibody domains selected using phage, it is

particularly important for the selection of Fv fragments which are formed by the non-covalent association of VH and VL fragments. Fv fragments have a tendency to dissociate and have a much reduced half-life in 30 circulation compared to whole antibodies. Fy fragments are displayed on the surface of phage, by the association of one chain expressed as a gene III protein fusion with the complementary chain expressed as a soluble fragment. If pairs of chains have a high tendency to dissociate, they will be much less likely to be selected as pabs. 35

Therefore, the population will be enriched for pairs which do associate stably. Although dissociation is less of a problem with Fab fragments, selection would also occur for Fab fragments which associate stably. pAbs allow selection for stability to protease attack, only those pAbs that are not cleaved by protesses will be capable of binding their ligand and therefore populations of phage will be enriched for those displaying stable

antibody domains.

The technique of displaying binding molecules on the phage surface can also be used as a primary cloning system. For example, a cDNA library can be constructed and inserted into the bacteriophage and this phage library screened for the ability to bind a ligand. The ligand/binding molecule combination could include any pair of molecules with an ability to specifically bind to

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one another e.g. receptor/ligand, enzyme/substrate (or analogue), nucleic acid binding protein/nucleic acid etc. If one member of the complementary pair is available, this may be a preferred way of isolating a clone for the

5 other member of the pair.

It will often be necessary to increase the diversity of a population of genes cloned for the display of their proteins on phage or to mutate an individual nucleotide sequence. Although in vitro or in vivo mutagenesis

techniques could be used for either purpose, a particularly suitable method would be to use mutator strains. A mutator strain is a strain which contains a genetic defect which causes DNA replicated within it to be mutated with respect to its parent DNA. Hence if a population of genes as gene III fusions is introduced

population of genes as gene III fusions is introduced into these strains it will be further diversified and can then be transferred to a non-mutator strain, if desired, for display and selection. Example 38 covers the use of mutator strains with phage antibodies (an example of in vitro mutagenesis and selection of phage antibodies is

given in example 45).

Targeted gene transfer

A useful and novel set of applications makes use of the binding protein on the phage to target the phage 25 genome to a particular cell or group of cells. For example, a pab specific for a cell surface molecule could

be used to bind to the target cell via the surface molecule. The phage could then be internalised, either through the action of the receptor itself or as the result of another event (e.g. an electrical discharge such as in the technique of electroporation). The phage

such as in the technique of electroporation). The phage genome would then be expressed if the relevant control signals (for transcription and translation and possibly replication) were present. This would be particularly useful if the phage genome contained a sequence whose

expression was desired in the target cell (along with the appropriate expression control sequences). A useful sequence might confer antibiotic resistance to the recipient cell or label the cell by the expression of its product (e.g. if the sequence expressed a detectable gene

product such as a luciferase, see White, M, et al, Techniques 2(4), p194-201 (1990)), or confer a particular property on the target cell (e.g. if the target cell was a tumour cell and the new sequence directed the

45 expression of a tumour suppressing gene), or express an antisense construct designed to turn off a gene or set of genes in the target cell, or a gene or gene product designed to be toxic to the target cell.

Alternatively, the sequence whose expression is desired in the target cell can be encoded on a phagemid. The phagemid DNA may then be incorporated into a phage displaying an antibody specific for a cell surface

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receptor. For example, incorporation may be by superinfection of bacteria containing the phagemid, with a helper phage whose genome encodes the antibody fragment specific for the target cell. The package is then used to direct the phagemid to the target cell.

This technique of "targeted gene transfer" has a number of uses in research and also in therapy and diagnostics. For example, gene therapy often aims to target the replacement gene to a specific cell type that is deficient in its activity. Targetting pAbs provide a

means of achieving this.

In diagnostics, phage specific for particular bacteria or groups of bacteria have been used to target marker genes, e.g. luciferase, to the bacterial host 15 (sec. for example, Ulitzer, S., and Kuhn, J., EPA 85303913.9). If the host range of the phage is appropriate, only those bacteria that are being tested for, will be infected by the phage, express the luciferase gene and be detected by the light they emit.

This system has been used to detect the presence of 20 Salmonella. One major problem with this approach is the initial isolation of a bacteriophage with the correct host range and then the cloning of a luciferase gene cassette into that phage, such that it is functional. The pAb system allows the luciferase cassette to be

cloned into a well characterised system (filamentous phage) and allows simple selection of an appropriate host range, by modifying the antibody (or other binding molecule) specificity that the pab encodes. 30

The present applicants have also been able to develop novel selection systems and assay formats which depend on the unique properties of these replicable

genetic display packages e.g. pAbs. TERMINOLOGY

35 Much of the terminology discussed in this section has been mentioned in the text where appropriate. Specific Binding Pair

This describes a pair of molecules (each being a member of a specific binding pair) which are naturally derived or synthetically produced. One of the pair of molecules, has an area on its surface, or a cavity which specifically binds to, and is therefore defined as complementary with a particular spatial and polar organisation of the other molecule, so that the pair have the property of binding specifically to each other.

Examples of types of specific binding pairs are antigenantibody. biotin-avidin, hormone-hormone receptor, receptor-ligand, enzyme-substrate, lgG-protein A.

Multimeric Member

This describes a first polypeptide which will associate with at least a second polypeptide, when the polypeptides are expressed in free form and/or on the surface of a substrate. The substrate may be provided by a bacteriophage. Where there are two associated polypeptides, the associated polypeptide complex is a dimer, where there are three, a trimer etc. The dimer, trimer, multimer etc or the multimeric member may

comprise a member of a specific binding pair.

Example multimeric members are heavy domains based on an immunoglobulin molecule, light domains based on an immunoglobulin molecule. T-cell receptor subunits.

10 Replicable Genetic Display Package (Rgdp)

This describes a biological particle which has genetic information providing the particle with the ability to replicate. The particle can display on its surface at least part of a polypeptide. The polypeptide in the particle and/or artificially placed into the particle or an ancestor of it. The displayed polypeptide may be any member of a specific binding pair eg. heavy or light chain domains based on an immunoglobulin molecule, an enzyme or a receptor etc.

The particle may be a virus eg. a bacteriophage such

as fd or M13.

Package
This describes a replicable genetic display package
in which the particle is displaying a member of a
specific binding pair at its surface. The package may be
a bacteriophage which displays an antigen binding domain
at its surface. This type of package has been called a

phage antibody (pAb). 30 Antibody

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Antibody
This describes an immunoglobulin whether natural or
partly or wholly synthetically produced. The term also
covers any protein having a binding domain which is
homologous to an immunoglobulin binding domain. These
proteins can be derived from natural sources, or partly
or wholly synthetically produced.

Example antibodies are the immunoglobulin isotypes and the Fab. F(ab¹)₂, scFv, Fv, dAb, Fd fragments.

Immunoglobulin Superfamily

This describes a family of polypeptides, the members of which have at least one domain with a structure related to that of the variable or constant domain of immunoglobulin molecules. The domain contains two β-sheets and usually a conserved disulphide bond (see A.F. Williams and A.N. Barclay 1988 Ann. Rev Immunol. 6

381-405).
Example members of an immunoglobulin superfamily are

CD4. platelet derived growth factor receptor (PDGFR), intercellular adhesion molecule. (ICAM). Except where the context otherwise dictates, reference to immunoglobulins and immunoglobulin homologs in this application includes members of the immunoglobulin

superfamily and homologs thereof. Homologs

This term indicates polypeptides having the same or conserved residues at a corresponding position in their primary, secondary or tertiary structure. The term also extends to two or more nucleotide sequences encoding the homologous polypeptides.

Example homologous peptides are the immunoglobulin isotypes.

10 Functional

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In relation to a sbp member displayed on the surface of a rgdp, means that the sbp member is presented in a folded form in which its specific binding domain for its complementary sbp member is the same or closely analogous to its native configuration, whereby it exhibits similar specificity with respect to the complementary sbp member.

In this respect, it differs from the peptides of Smith et al, supra, which do not have a definite folded configuration and can assume a variety of configurations of determined by the complementary members with which they

determined by the complementary members with which they may be contacted.

Genetically diverse population

In connection with sbp members or polypeptide components thereof, this is referring not only to diversity that can exist in the natural population of cells or organisms, but also diversity that can be created by artificial mutation in vitro or in vivo.

Mutation in vitro may for example, involve random mutagenesis using Oligonucleotides having random mutations of the sequence desired to be varied. In vivo mutagenesis may for example, use mutator strains of host microorganisms to harbour the DNA (see Example 38 below).

Domain

A domain is a part of a protein that is folded
within itself and independently of other parts of the
same protein and independently of a complementary binding
member.

Folded Unit

This is a specific combination of an α -helix and/or β -strand and/or β -turn structure. Domains and folded units contain structures that bring together amino acids that are not adjacent in the primary structure. Free Form

This describes the state of a polypeptide which is not displayed by a replicable genetic display package.

Conditionally Defective

This describes a gene which does not express a particular polypeptide under one set of conditions, but expresses it under another set of conditions. An example, is a gene containing an amber mutation expressed in non-suppressing or suppressing hosts respectively.

Alternatively, a gene may express a protein which is

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defective under one set of conditions, but not under An example is a gene with a temperature another set. sensitive mutation.

Suppressible Translational Stop Codon

This describes a codon which allows the translation 5 of nucleotide sequences downstream of the codon under one set of conditions, but under another set of conditions translation ends at the codon. Example of suppressible translational stop codons are the amber, ochre and opal codons. 10

Mutator Strain

This is a host cell which has a genetic defect which causes DNA replicated within it to be mutated with respect to its parent DNA. Example mutator strains are

NR9046mutD5 and NR9046 mut T1 (see Example 38). 15

This is a phage which is used to infact cells containing a defective phage genome and which functions to complement the defect. The defective phage genome can be a phagemid or a phage with some function encoding gene sequences removed. Examples of helper phages are M13K07, MIGKO7 cene III no. 3; and phage displaying or encoding a binding molecule fused to a capsid protein. Vector

This is a DNA molecule, capable of replication in a 25 host organism, into which a gene is inserted to construct a recombinant DNA molecule.

Phage Vector

This is a vector derived by modification of a phage genome, containing an origin of replication for a 30 bacteriophage, but not one for a plasmid. Phagemid Vector

This is a vector derived by modification of a plasmid genome, containing an origin of replication for a bacteriophage as well as the plasmid origin of replication.

Secreted

This describes a rgdp or molecule that associates with the member of a sbp displayed on the rgdp, in which the sbp member and/or the molecule, have been folded and 40 the package assembled externally to the cellular cytosol. Repertoire of Rearranged Immunoglobulin Genes

A collection of naturally occurring nucleotides eg DNA sequences which encoded expressed immunoglobulin genes in an animal. The sequences are generated by the 45 in vivo rearrangement of eg V, D and J segments for H chains and eg the V and J segments for L chains. Alternatively the sequences may be generated from a cell line immunised in vitro and in which the rearrangement in response to immunisation occurs intracellularly. 50

Library

A collection of nucleotide eg DNA, sequences within

clones.

Repertoire of Artificially Rearranged Immunoglobulin

A collection of nucleotide eg DNA, sequences derived wholly or partly from a source other than the rearranged immunoglobulin sequences from an animal. include for example. DNA sequences encoding VH domains by combining unrearranged V segments with D and J segments and DNA sequences encoding VL domains by combining V and 10 J segments.

Part or all of the DNA sequences may be derived by oligonucleotide synthesis.

Secretory Leader Peptide

This is a sequence of amino acids joined to the N-15 terminal end of a polypeptide and which directs movement of the polypeptide out of the cytosol. Eluant

This is a solution used to breakdown the linkage between two molecules. The linkage can be a non-covalent 20 or covalent bond(s). The two molecules can be members of a sbp.

Derivative

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This is a substance which derived from a polypeptide which is encoded by the DNA within a selected roop. The 25

derivative polypeptide may differ from the encoded polypeptide by the addition, deletion, substitution or insertion of amino acids, or by the linkage of other molecules to the encoded polypetide. These changes may be made at the nucleotide or protein level. For example 30 the encoded polypeptide may be a Fab fragment which is

then linked to an Fc tail from another source. Alternatively markers such as enzymes, flouresceins etc may be linked to eg Fab, scFv fragments.

The present invention provides a method for 35 producing a replicable genetic display package or population such rgdps of which method comprises the steps of:

a) inserting a nucleotide sequence encoding a member of a specific binding pair eg. a binding molecule within a viral genome;

bì culturing the virus containing said nucleotide sequence, so that said binding molecule is expressed and displayed by the virus at its surface.

The present invention also provides a method for 45 selecting a rgdp specific for a particular epitope which comprises producing a population of such roops as described above and the additional step of selecting for said binding molecule by contacting the population with said epitope so that individual rodps with the desired specificity may bind to said epitope. The method may comprise one or more of the additional steps of: (i) separating any bound rgdps from the epitope; (ii) 50

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recovering any separated rgdps and (iii) using the inserted nucleotide sequences from any separated rgdps in a recombinant system to produce the binding molecule separate from virus. The selection step may isolate the nucleotide sequence encoding the binding molecule of desired specificity, by virtue of said binding molecule being expressed in association with the surface of the virus in which said encoding nucleic acid is contained.

The present invention also provides a method of 10 producing a multimeric member of a specific binding pair

(sbp), which method comprises:

expressing in a recombinant host organism a first polypeptide chain of said sbp member or a genetically diverse population of said sbp member fused to a component of a secreted replicable genetic display package (rgdp) which thereby displays said polypeptide at the surface of the package, and expressing in a recombinant host organism a second polypeptide chain of said multimer and causing or allowing the polypeptide chains come together to form said multimer as part of said rgdp at least one of said polypeptide chains being expressed from nucleic acid that is capable of being packaged using said component therefor, whereby the genetic material of each said rgdp encodes a said

25 polypeptide chain. Both said chains may be expressed in the same host organism.

The first and second chains of said multimer may be expressed as separate chains from a single vector 30 containing their respective nucleic acid.

At least one of said polypeptide chains may be

expressed from a phage vector.

At least one of said polypeptide chains may be expressed from a phagemid vector, the method including using a helper phage, or a plasmid expressing complementing phage genes, to help package said phagemid genome, and said component of the rgdp is a capsid protein therefor. The capsid protein may be absent, defective or conditionally defective in the helper phage.

The method may comprise introducing a vector capable of expressing said first polypeptide chain, into a host organism which expresses said second polypeptide chain in free form, or introducing a vector capable of expressing said second polypeptide in free form into a host organism

45 which expresses said first polypeptide chain.

Each of the polypeptide chain may be expressed from nucleic acid which is capable of being packaged as a rgdp using said component fusion product, whereby encoding nucleic acid for both said polypeptide chains are packaged in respective rgdps.

The nucleic acid encoding at least one of said first and second polypeptide chains may be obtained from a

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thereof.

library of nucleic acid including nucleic acid encoding said chain or a population of variants of said chain. Both the first and second polypeptide chains may be obtained from respective said libraries of nucleic acid.

The present invention also provides a method of producing a member of a specific binding pair (sbp), from a nucleic acid library including nucleic acid encoding said sbp member or a genetically diverse population of said type of sbb members, which method comprises:

expressing in recombinant host calls polypeptides encoded by said library nucleic acid fused to a component of a secreted replicable genetic display package (rgdp) or in free form for association with a polypeptide component of said spp member which is expressed as a fusion to said rgdp component so that the rgdp displays said sbp member in functional form at the surface of the package, said library nucleic acid being contained within the host cells in a form that is capable of being packaged using said rgdp component, whereby the genetic material of an rgdp displaying an sbp member contains nucleic acid encoding said sbp member or a polypeptide component

The nucleotide sequences for the libraries may be derived from eg animal spleen cells or peripheral blood lymphocytes. Alternatively the nucleotide sequence may be derived by the in <u>vitro</u> mutagenesis of an existing antibody coding sequence.

The present invention also provides a method of producing a member of a specific binding pair (sbp), which method comprises:

expressing in recombinant host cells nucleic acid encoding said sbp member or a genetically diverse population of said type of sbp member wherein the or each said sbp member or a polypeptide component thereof is expressed as a fusion with a component of a secreted replicable genetic display package (rgdp) which displays said sbp member at the surface of the package, nucleic acid encoding said sbp member or a polypeptide component thereof being contained within the host cell in a form that is capable of being packaged using said rgdp component whereby the genetic material of the rgdp displaying said sbp member encodes said sbp member or a polypeptide component thereof, said host organism being a mutator strain which introduces genetic diversity into the sbp member to produce said mixed population.

The present invention also provides a method of producing a member of a specific binding pair (sbp), which method comprises:

expressing in recombinant host cells nucleic acid

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encoding said sbp member or a genetically diverse population of said type of sbp member wherein the or each said sbp member or a polypeptide component thereof is expressed as a fusion with a component of a secreted replicable genetic display package (rgdp) which displays said sbp member in functional form at the surface of the package, nucleic acid encoding said sbp member or a polypeptide component thereof being contained within the host cell in a form that is capable of being packaged using said rgdp component whereby the genetic material of the rgdp displaying an sbp member encodes said sbp member or a polypeptide component thereof, said fusions being with bacteriophage capsid protein and the rgdps being formed with said fusions in the absence of said capsid expressed in wild-type form. The present invention also provides a method of

The present invention also provides a method of producing a member of a specific binding pair (sbp) which

method comprises:

expressing in recombinant host cells nucleic acid encoding said sbp member or a genetically diverse population of said type of sbp member or a polypeptide component thereof fused to a component of a secreted replicable genetic display package (rgdp) which displays said sbp member in functional form at the surface of the package, nucleic acid encoding said sbp member or a polypeptide component thereof being contained within the host cell in a form that is capable of being packaged using said rgdp component whereby the genetic material of the rgdp displaying an sbp member or a polypeptide component thereof encodes said sbp member or a polypeptide component thereof, said sbp member or polypeptide component thereof being expressed from a phagemid as a capsid fusion, and a helper phage, or a plasmid expressing complementing phage genes, is used along with said capsid fusions to package the phagemid nucleic acid.

The library or genetically diverse population may be

40 obtained from:

(i) the repertoire of rearranged immunoglobulin genes of an animal immunised with complementary sbp member,

(ii) the repertoire of rearranged immunoglobulin genes of an animal not immunised with complementary sbo member.

(iii) a repertoire of artificially rearranged immunoglobulin gene or genes

(v) a mixture of any of (i), (ii), (iii) and (iv). The capsid protein may be absent, defective or

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conditionally defective in the helper phage.

The host cell may be a mutator strain which introduces genetic diversity into the sbp member nucleic acid.

S The sbp member may comprise a domain which is, or is homologous to, an immunoglobulin domain.

The rgdp may be a bacteriophage, the host a bacterium, and said component of the rgdp a capsid protein for the bacterophage. The phage may be a filamentous phage. The phage may be selected from the class I phages fd, M13, fl. Ifl, lke, ZJ/Z, Ff and the class II phages Xf, Pfl and Pf3. The phage may be fd or a derivative of fd. The derivative may be tetracycline resistant. The said sbp member or polypeptide chain

thereof may be expressed as a fusion with the gene III capsid protein of phage fd or its counterpart in another filementous phage. The sbp member or polypeptide chain thereof may be inserted in the N-terminal region of the mature capsid protein downstream of a secretory leader

20 peptide. The sequence may be inserted after amino acid +1 of the mature protein. The site for insertion may be flanked by short sequences corresponding to sequences which occur at each end of the nucleic acid to be inserted. For example where 4 the protein domain is an

25 immunoglobulin domain, the insertion site in the phage may be flanked by nucleotide sequences which code for the first five amino acids and the last five amino acids of the Ig domain. Such flanking nucleotide sequences are shown in figure 4(2) B and C, wherein the site-flanking

nucleotide sequences encode amino acid sequences QVQLQ 30 and VTVSS which occur at either end of the VH domain, or QVQLQ and LEIKR which occur at either end of the Fv (combined VH + VL) domain. Each of these sequences flanking the insertion site may include a suitable cleavage site, as shown in Fig 4.

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Alternatively, the flanking nucleotide sequences shown in figure 4(2)B and C as described above, may be used to flank the insertion site for any nucleic acid to be inserted, whether or not that nucleic acid codes an immunoclobulin.

The host may be E.coli.

Nucleic acid encoding an sbp member polypeptide may be linked downstream to a viral capsid protein through a suppressible translational stop codon.

As previously mentioned, the present invention also provides novel selection systems and assay formats. these systems and formats, the gene sequence encoding the binding molecule (eg. the antibody) of desired specificity is separated from a general population of rgdps having a range of specifies, by the fact of its binding to a specific target (eg the antigen or epitope). Thus the rgdps formed by said expression may be selected

or screened to provide an individual sbp member or a selected mixed population of said sbp members associated in their respective rgdps with nucleic acid encoding said sbp member or a polypeptide chain thereof. The rgdps may be selected by affinity with a member complementary to

said sbp member.

Any rgdps bound to said second member may be recovered by washing with an eluant. The washing conditions may be varied in order to obtain rodps with different binding affinities for said epitope. Alternatively, to obtain eg high affinity rgdps, the 1.0 complementary member (eg an epitope) may be presented to the population of rgdps (eg pAbs) already bound to a binding member in which case pabs with a higher affinity for the epitope will displace the already bound binding 15 member. Thus the eluant may contain a molecule which competes with said rgdp for binding to the complementary sbp member. The rgdp may be applied to said complementary sbp member in the presence of a molecule which competes with said package for binding to said 20 complementary sbp member. Nucleic acid derived from a selected or screened rgdp may be used to express said sbp member or a fragment or derivative thereof in a recombinant host organism. Nucleic acid from one or more rgdps may be taken and used to provide encoding nucleic 25 acid in a further said method to obtain an individual sbp member or a mixed population of sbp members, or encoding nucleic acid therefor. The expression end product may be modified to produce a derivative thereof.

The expression end product or derivative thereof may be used to prepare a therapeutic or prophylactic

medicament or a diagnestic product.

The present invention also provides recombinant host cells harbouring a library of nucleic acid fragments comprising fragments encoding a genetically diverse population of a type of member of a specific binding pair (sbp), each sbp member or a polypeptide component thereof being expressed as a fusion with a component of a secretable replicable genetic display package (rgdp), so that said sbp members are displayed on the surface of the 40 rgdps in functional form and the genetic material of the rodps encode the associated sbp member or a polypeptide The type of sbp members may be component thereof. immunoglobulins or immunoglobulin homologs, a first

polypeptide chain of which is expressed as a said fusion 45 with a component of the rgdp and a second polypeptide chain of which is expressed in free form and associates with the fused first polypeptide chain in the rgdp. The present invention also provides a helper phage

whose genome lacks nucleic acid encoding one of its :-0 capsid proteins, or whose encoding nucleic acid therefor is conditionally defective, or which encodes said capsid

protein in defective or conditionally defective form.

The present invention also provides a bacterial host cell containing a filamentous phage genome defective for a capsid protein thereof and wherein the host cell is capable of expressing capsid protein complementing said defect such that infectious phage particles can be obtained therefrom. The complementing capsid protein may be expressed in said host from another vector contained therein. The defective capsid protein may be gene III of phage fd or its counterpart in another filamentous phage.

The present invention also provides recombinant E.coli TG1 M13K07 gIII No. 3 (NCTC 12478).

The present invention also provides a phage antibody having the form of a replicable genetic display package 15 displaying on its surface in functional form a member of a specific binding pair or a specific binding domain thereof.

In the above methods, the binding molecule may be an antibody, or a domain that is homologous to an immunoglobulin. The antibody and/or domain may be either naturally derived or synthetic or a combination of both. The domain may be a Fab, scFv, Fv dAb or Fd molecule. Alternatively, the binding molecule may be an enzyme or receptor or fragment, derivative or analogue of any such enzyme or receptor. Alternatively, the binding molecule may be a member of an immunoglobulin superfamily and

which has a structural form based on an immunoglobulin molecule.

The present invention also provides rgdps as defined above and members of specific binding pairs eg. binding

molecules such as antibodies, enzymes, receptors, fragments and derivatives thereof, obtainable by use of any of the above defined methods. The derivatives may comprise members of the specific binding pairs fused to another molecule such as an enzyme or a Fc tail.

The invention also includes kits for carrying out the methods hereof. The kits will include the necessary vectors. One such vector will typically have an origin of replication for single stranded bacteriophage and either contain the sbp member nucleic acid or have a restriction site for its insertion in the 5' end region

restriction site for its insertion in the 5' end region of the mature coding sequence of a phage capsid protein, and with a secretory leader coding sequence upstream of said site which directs a fusion of the capsid protein exogenous polypeptide to the periplasmic space.

The restriction sites in the vectors are preferably those of enzymes which cut only rarely in protein coding sequences.

The kit preferably includes a phagemid vector which may have the above characteristics, or may contain, or have a site for insertion, of sbp member nucleic acid for expression of the encoded polypeptide in free form.

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The kits will also contain ancillary components required for carrying out the method, the nature of such components depending of course on the particular method employed.

5 Useful ancillary components may comprise helper phage, PCR primers, and buffers and enzymes of various kinds.

PCR primers and associated reagents for use where the sbp members are antibodies may have the following characteristics:

(i) primers having homology to the 5' end of the sense or anti-sense strand of sequences encoding domains of antibodies; and

(ii) primers including tag sequences 5' to these homologous sequences which incorporate restriction sites to allow insertion into vectors; together with sequences to allow assembly of amplified VH and VL regions to enable expression as Fv. scFv or Fab framents.

20 Buffers and enzymes are typically used to enable preparation of nucleotide sequences encoding Fy, scFv or Fab fragments derived from rearranged or unrearranged immunoglobulin genes according to the strategies described herein.

25 The applicants have chosen the filamentous Fspecific bacteriophages as an example of the type of
phage which could provide a vehicle for the display of
binding molecules e.g. antibodies and antibody fragments
and derivatives thereof, on their surface and facilitate
30 subsequent selection and manipulation.

The F-specific phages (e.g. fl. fd and M13) have svolved a method of propagation which does not kill the host cell and they are used commonly as vehicles for recombinant DNA (Kornberg, A., DNA Replication, W.H. Freeman and Co., San Francisco, 1980). The single stranded DNA genome (approximately 6.4 kb) of fd is extruded through the bacterial membrane where it sequesters capsid sub-units, to produce mature virions. These virions are 6 nm in diameter, lym in length and each contain approximately 2,800 molecules of the major coat protein encoded by viral gene VIII and four molecules of the adsorption molecule gene III protein (g3p) the latter is located at one end of the virion. The structure has been reviewed by Webster at al., 1978

The structure has been reviewed by Webster et al., 1978 in The Single Stranded DNA Phages, 557-569, Cold Spring Harbor Laboratory Press. The gene III product is involved in the binding of the phage to the bacterial Fpilus.

Although these phages do not kill their host during normal replication, disruption of some of their genes can lead to cell death (Kornberg, A., 1980 supra.) This places some restraint on their use. The applicants have

of e.g. antibodies.

recognized that gene III of phage fd is an attractive possibility for the insertion of biologically active foreign sequences. There are however, other candidate sites including for example gene VIII and gene VI.

The protein itself is only a minor component of the phage coat and disruption of the gene does not lead to cell death (Smith, G. 1988, Virology 167: 156-165). Furthermore, it is possible to insert some foreign sequences (with no biological function) into various 10 positions within this dene (Smith, G. 1985 Science 228: 1315-1317., Parmley, S.F. and Smith, G.P. Gene: 73 (1988) p. 305-318., and de la Cruz, V.F., et al., 1988, J. Biol. Chem., 263: 4318-4322). Smith et al described the display of peptides on the outer surface of phage but 15 they did not describe the display of protein domains. Peptides can adopt a range of structures which can be different when in free solution, than when bound to, for example, an antibody, or when forming part of a protein (Stanfield, R.I. et al., (1990) Science 248, p712-719). 20 Proteins in general have a well defined tertiary structure and perform their biological function only when adopting this structure. For example, the structure of the antibody D1.3 has been solved in the free form and when bound to antigen (Bhat, T.N. et al., (1990) Nature 25 347, p483-4851. The gross structure of the protein is identical in each instance with only minor variations around the binding site for the antigen. Other proteins have more substantial conformation changes on binding of ligand, for instance the enzymes hexokinase and pyruvate 30 dehydrogenase during their catalytic cycle, but they still retain their overall pattern of folding. structural integrity is not confined to whole proteins. but is exhibited by protein domains. This leads to the concept of a folded unit which is part of a protein, 35 often a domain, which has a well defined primary, secondary and tertiary structure and which retains the same overall folding pattern whether binding to a binding partner or not. The only game sequence that Smith et al., described that was of sufficient size to encode a domain (a minimum of perhaps 50 amino acids) was a 335bp 40 fragment of a \$-galctrosidase corresponding to nucleotides 861-1195 in the 8-galactosidase gene sequence (Parmley, S. + Smith, G.P. 1988 supra. This would encode 112 amino acids of a much larger 380 amino acid domain. Therefore, prior to the present application, no 45 substantially complete domain or folded unit had been displayed on phage. In these cases, although the

sequences could be detected on the phage surface by use The protein encoded by gene III has several domains (Pratt, D., et al., 1969 Virology 39:42-53., Grant, R.A.,

infectivity of the virion was disrupted, the inserted

et al., 1981, J. Biol. Chem. 256: 539-546 and Armstrong, J., et al., FEBS Lett. 135: 167-172 1981.) including: (1) a signal sequence that directs the protein to the cell membrane and which is then cleaved off; (i1) a domain that anchors the mature protein into the bacterial cell membrane (and also the phage coat); and (iii) a domain that specifically binds to the phage receptor, the Fpilus of the host bacterium. Short sequences derived from protein molecules have been inserted into two places within the mature molecule (Smith, G., 1985 supra., and Parmley, S.F. and Smith G.P., 1988 supra.). Namely, into an inter-domain region and also between amino acids 2 and 3 at the N-terminus. The insertion sites at the N-

an inter-womain region and also between daths actus 2 and 3 at the N-terminus. The insertion sites at the N-terminus were more successful in maintaining the structural integrity of the gene III protein and displaying the peptides on the surface of the phage. By use of antisera specific for the peptides, the peptides inserted into this position were shown to be on the surface of the phage. These authors were also able to purify the phage, using this property. However, the peptides expressed by the phage, did not possess

measurable biological functions of their own.

Retaining the biological function of a molecule when it is expressed in a radically different context to its natural state is difficult. The demands on the structure 25 of the molecule are heavy. In contrast, retaining the ability to be bound by specific antisera is a passive process which imposes far less rigorous demands on the structure of the molecule. For example, it is the rule rather than the exception that polyclonal antisera will 30 recognise totally denatured, and biologically inactive, proteins on Western blots (see for example, Harlow, E. and Lane, D., Antibodies, a Laboratory Manual, Cold Spring Harbor Laboratory Press 1988). Therefore, the insertion of peptides into a region that allows their 35 structure to be probed with antisers teaches only that the region allows the inserted sequences to be exposed and does not teach that the region is suitable for the insertion of large sequences with demanding structural 40

constraints for the display of a molecule with a biological or binding function. In particular, it does not teach that domains or folded units of proteins can be displayed from sequences inserted in this region.

This experience with Western blots is a graphic practical demonstration which shows that retaining the ability to be bound by specific antisers imposes far less rigorous demands on the structure of a polypeptide, than does folding for the retention of a biological function. Studies have been carried out, in which E.coli have

50 been manipulated to express the protein B-adranargic receptor as a fusion with the outer membrane protein lam8. The B-adranargic receptor was expressed in a functional form as determined by the presence of binding activity. However, when an equivalent antibody fusion was made with lamB, the antibody fusion was toxic to the host cell.

The applicants have investigated the possibility of inserting the gene coding sequence for biologically active antibody fragments into the gene III region of fd to express a large fusion protein. As is apparent from the previous discussion, this approach makes onerous demands on the functionality of the fusion regions.

demands on the functionality of the fusion protein. The insertion is large, encoding antibody fragments of at least 100-200 amino acids; the antibody derived domain must fold efficiently and correctly to display antigenbinding; and most of the functions of gene III must be

15 retained. The applicants approach to the construction of the fusion molecule was designed to minimise the risk of disrupting these functions. In an embodiment of the invention, the initial vector used was fd-tet (Zacher, A.N., et al. 1980, Gene 9. 127-140) a tetracycline

20 resistant version of fd bacteriophage that can be propagated as a plasmid that confers tetracycline resistance to the infected <u>E.coli</u> host. The applicants chose to insert after the signal sequence of the fd gene III protein for several reasons. In particular, the

applicants chose to insert after amino acid 1 of the mature protein to retain the context for the signal peptidase cleavage. To retain the structure and function of gene III itself, the majority of the original amino acids are synthesized after the inserted immunoglobulin sequences. The inserted immunoglobulin sequences were designed to include the sequences were

designed to include residues from the switch region that links VH-VL to CHI-CL (Lesk, A., and Chothia, C., Nature 335, 188-190, 1988).

Surprisingly, by manipulating gene III of stateriophage fd, the present applicants have been able to construct a bacteriophage that displays on its surface large biologically functional antibody, enzyme, and receptor molecules whilst remaining intact and infectious. Furthermore, the phages bearing antibodies of desired specificity, can be selected from a background

of phages not showing this specificity.

The sequences coding for a population of antibody molecules and for insertion into the vector to give expression of antibody binding functions on the phage surface can be derived from a variety of sources. For example, immunised or non-immunised rodents or humans, and from organs such as spleen and peripheral blood lymphocytes. The coding sequences are derived from these sources by techniques familiar to those skilled in the art (Orlandi, R., et al., 1989 supra; Larrick, J.W., et al., 1989 supra; Chiang, Y.L., et al., 1989 supra; Techniques Z, p. 360-366; Ward, E.S, et al., 1989 supra;

Sastry, L., et al., 1989 supra.) or by novel linkage strategies described in examples 14, 33, 40 and 42. Novel strategies are described in examples 7, 25, 33, 39 and 40 for displaying dimeric molecules eg Fab and Fv fragments on the surface of a phage. Each individual pab in the resulting library of pabs will express antibodies.

in the resulting library of pAbs will express antibodies or antibody derived fragments that are monoclonal with respect to their antigen-binding characteristics.

The disclosure made by the present applicants is

The disclosure made by the present approach is important and provides a significant breakthrough in the technology relating to the production of biological binding molecules, their fragments and derivatives by the use of recombinant methods.

In standard recombinant techniques for the production of antibodies, an expression vector containing sequences coding for the antibody polypeptide chains is used to transform e.g. E.coli. The antibody polypeptides are expressed and detected by use of standard screening systems. When the screen detects an antibody polypeptide of the desired specificity, one has to return to the particular transformed E.coli expressing the desired antibody polypeptide. Furthermore, the vector containing the coding sequence for the desired antibody polypeptide then has to be isolated for use from E.coli in further

25 processing steps.

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In the present invention however, the desired antibody polypeptide when expressed, is already packaged with its gene coding sequence. This means that when the an antibody polypeptide of desired specificity is selected, there is no need to return to the original culture for isolation of that sequence. Furthermore, in previous methods in standard recombinant techniques, each clone expressing antibody needs to be screened individually. The present application provides for the selection of clones expressing antibodies with desired properties and thus only requires screening of clones from an enriched pool.

Because a rgdp (eg a pAb) is a novel structure that displays a member of a specific binding pair (eg an antibody of monoclonal antigen-binding specificity) at the surface of a relatively simple replicable structure also containing the genetic information encoding the member, rgdps eg pAbs, that bind to the complementary member of the specific binding pair (eg antigen) can be recovered very efficiently by either eluting off the complementary member using for example diethylamine, high salt etc and infecting suitable bacteria, or by denaturing the structure, and specifically amplifying the sequences encoding the member using FCR. That is, there is no necessity to refer back to the original bacterial

clone that gave rise to the pAb.

For some purposes, for example immunoprecipitation

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and some diagnostic tests, it is advantageous to use polyclonal antibodies or antibody fragments. The present invention allows this to be achieved by either selection of an enriched pool of pAbs with desired properties or by mixing individually isolated clones with desired The antibodies or antibody fragments may properties. then be expressed in soluble form if desired. Such a selected polyclonal pAb population can be grown from stocks of phage, bacteria containing phagemids or 10 bacteria expressing soluble fragments derived from the selected polyclonal population. Thus a readent equivalent to a polyclonal antiserum is created which can

be replicated and routinely manufactured in culture without use of animals. SELECTION FORMATS AND AFFINITY MATURATION

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Individual rgdps eg pAbs expressing the desired specificity eg for an antigen, can be isolated from the complex library using the conventional screening

techniques (e.g. as described in Harlow, E., and Lane, 20 D., 1988, supra Gherardi, E et al. 1990. J. Immunol.

meth, 126 p61-68).

The applicants have also devised a series of novel selection techniques that are practicable only because of the unique properties of rgdps. The general outline of 25 some screening procedures is illustrated in figure 2 using pabs as an example type of rodp.

The population/library of pAbs to be screened could be generated from immunised or other animals; or be created in vitro by mutagenising pre-existing phage antibodies (using techniques well-known in the art such as oligonuclectide directed mutagenesis (Sambrook, J., et al., 1989 Molecular Cloning a Laboratory Manual, Cold Spring Harbor Laboratory Press). This population can be screened in one or more of the formats described below with reference to figure 2, to derive those individual pAbs whose antigen binding properties are different from sample c.

Binding Elution

soak solution, etc.

Figure 2(i) shows antigen (ag) bound to a solid 40 surface (s) the solid surface (s) may be provided by a petri dish, chromatography beads, magnetic beads and the like. The population/library of pAbs is then passed over the ag, and those individuals p that bind are retained after washing, and optionally detected with detection 45 system d. A detection system based upon anti-fd antisera is illustrated in more detail below in example 4. samples of bound population p are removed under increasingly stringent conditions, the binding affinity represented in each sample will increase. Conditions of increased stringency can be obtained, for example, by 50 increasing the time of soaking or changing the pH of the

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Competition

Referring to figure 2(ii) antigen ag can be bound to a solid support s and bound to saturation by the original binding molecule c. If a population of mutant pAb (or a set of unrelated pAbs) is offered to the complex, only those that have higher affinity for antigen ag than c will bind. In most examples, only a minority of population c will be displaced by individuals from population p. If c is a traditional antibody molecule, all bound material can be recovered and bound p recovered by infecting suitable bacteria and/or by use of standard

techniques such as PCR. An advantageous application is where ag is used as a receptor and c the corresponding ligand. The recovered bound population p is then related structurally to the receptor binding site/and or ligand. This type of specificity is known to be very useful in the

pharmaceutical industry.

Another advantageous application is where ag is an antibody and c its antigen. The recovered bound 20 population p is then an anti-idiotype antibody which have numerous uses in research and the diagnostic and pharmaceutical industries.

At present it is difficult to select directly for anti-idiotype antibodies. pabs would give the ability to 25 do this directly by binding pAb libraries (eg a naive library) to B cells (which express antibodies on their surface) and isolating those phage that bound well.

In some instances it may prove advantageous to preselect population p. For example, in the anti-idiotype example above, p can be absorbed against a related

antibody that does not bind the antigen. However, if c is a pAb, then either or both c and p can advantageously be marked in some way to both distinguish and select for bound p over bound c. This marking can be physical, for example, by pre-labelling p with biotin; or more advantageously, genetic. example, c can be marked with an EcoB restriction site, whilst p can be marked with an EcoK restriction site (see Carter, P. et al., 1985, Nucl. Acids Res. 13, 4431-4443). When bound p+c are eluted from the antigen and used to infect suitable bacteria, there is restriction (and thus no growth) of population c (i.e. EcoB restricting bacteria in this example). Any phage that grew, would be greatly enriched for those individuals from p with higher binding affinities. Alternatively, the genetic marking can be achieved by marking p with new sequences, which

can be used to specifically amplify p from the mixture using PCR. Since the bound pAbs can be amplified using for example PCR or bacterial infection, it is also possible to rescue the desired specificity even when insufficient individuals are bound to allow detection via conventional techniques.

The preferred method for selection of a phage displaying a protein molecule with a desired specificity or affinity wil. Often be elution from an affinity matrix with a ligand (eg example 21). Elution with increasing concentrations of ligand should elute phage displaying binding molecules of increasing affinity. However, when eg a pAb binds to its antigen with high affinity or 10 avidity (or another protein to its binding partner) it may not be possible to elute the pAb from an affinity matrix with molecule related to the antigen. Alternatively, there may be no suitable specific eluting molecule that can be prepared in sufficiently high 15 concentration. In these cases it is necessary to use an elution method which is not specific to eg the antigenantibody complex. Some of the non-specific elution methods generally used reduce phage viability for instance, phage viability is reduced with time at pH12 (Rossomando, E.F. and Zinder N.D. J. Mol.Biol. 36 387-399 20 1968). There may be interactions between eg antibodies and affinity matrices which cannot be disrupted without completely removing phage infectivity. In these cases a method is required to elute phage which does not rely on disruption of eg the antibody - antigen interaction. A method was therefore devised which allows elution of 25 bound pAbs under mild conditions (reduction of a dithiol

group with dithicthreifol) which do not disrupt phage structure (example 47).

This elution procedure is just one example of an elution procedure under mild conditions. A particularly advantageous method would be to introduce a nucleotide

sequence encoding amino acids constituting a recognition

site for cleavage by a highly specific protease between the foreign gene inserted, in this instance a gene for an antibody fragment, and the sequence of the remainder of gene III. Examples of such highly specific proteases are Factor X and thrombin. After binding of the phage to an affinity matrix and elution to remove non-specific

40 binding phage and weak binding phage, the strongly bound phage would be removed by washing the column with processe under conditions suitable for digestion at the cleavage site. This would cleave the antibody fragment from the phage particle eluting the phage. These phage

45 would be expected to be infective, since the only protease site should be the one specifically introduced. Strongly binding phage could then be recovered by infecting eg. E.coli TG1 cells.

An alternative procedure to the above is to take the affinity matrix which has retained the strongly bound pAb and extract the DNA, for example by boiling in SDS solution. Extracted DNA can then be used to directly

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transform E.coli host cells or alternatively the antibody encoding sequences can be amplified, for example using PCR with suitable primers such as those disclosed herein, and then inserted into a vector for expression as a soluble antibody for further study or a pAb for further rounds of selection.

Another preferred method for selection according to affinity would be by binding to an affinity matrix

containing low amounts of ligand.

If one wishes to select from a population of phages displaying a protein molecule with a high affinity for its ligand, a preferred strategy is to bind a population of phage to an affinity matrix which contains a low amount of ligand. There is competition between phage, displaying high affinity and low affinity proteins, for binding to the ligand on the matrix. Phage displaying high affinity protein is preferentially bound and low affinity protein is washed away. The high affinity protein is then recovered by elution with the ligand or by other procedures which elute the phage from the

affinity matrix (example 35 demonstrates this procedure). In summary then, for recovery of the packaged DN from the affinity step, the package can be simply eluted, it can be eluted in the presence of a homologous sbp member which competes with said package for binding to a

complementary sbp member; it could be removed by boiling, it could be removed by proteolytic clsevage of the protein; and other methods will be apparent to those skilled in the art eg. destroying the link between the substrate and complementary sbp member to release said packaged DNA and sbp member. At any rate, the objective is to obtain the DNA from the package so that it can be

used directly or indirectly, to express the sbp member encoded thereby.

The efficiency of this selection procedure for pabs and the ability to create very large libraries means that the immunisation techniques developed to increase the proportion of screened cells producing antibodies of interest will not be an absolute requirement. The technique allows the rapid isolation of binding specificities en actionable by those that would be difficult or even unobtainable by

conventional techniques, for example, catalytic or antiidiotypic antibodies. Removal of the animal altogether is now possible, once a complete library of the immune

repertoire has been constructed.

The novel structure of the pAb molecule can be used in a number of other applications, some examples of which are: Signal Amplification

Acting as a novel molecular entity in itself, rgdps eg pAbs combine the ability to bind a specific molecula eg antigen with amplification. if the major coat protein

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is used to attach another moiety. This moiety can be attached via immunological, chemical, or any other means and can be used, for example, to label the complex with detection reagents or cytotoxic molecules for use in vivo or in vitro.

Physical Detection

The size of the rgdps eg pAbs can be used as a marker particularly with respect to physical methods of detection such as electron microscopy and/or some biosensors, e.g. surface plasmon resonance. Diagnostic Assays

The rgdps eg pAbs also have advantageous uses in diagnostic assays, particularly where separation can be effected using their physical properties for example centrifugation, filtration etc.

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In order that the invention is more fully understood, embodiments will now be described in more detail by way of example only and not by way of limitation with reference to the figures described below. Figure 1 shows the basic structure of the simplest

antibody molecule IgG.

Figure 2 shows schematically selection techniques which utilise the unique properties of pAbs; 21) shows a binding/elution system: and (211) shows a competition system (p-pAb: ag-antigen to which binding by pAb is required; c-competitor population e.g. antibody, pAb, ligand; s-substrate (e.g. plastic beads etc); d-detection system.

Figure 3 shows the vector fd-tet and a scheme for the construction of vectors, fdTPs/8s (for insertion of 30 VH coding sequences) and fdTPs/Xh for the insertion of

scFv coding sequences.

Figure 4 shows the nucleotide sequences for the oligonucleotides and vectors. All sequences are drawn 5' 35 to 3' and are numbered according to Beck et al., 1978. Nucl. Acid Res., 5: 4495-4503. 4.1 shows the sequences of the oligonucleotides used for mutagenesis (oligo's 1 and 2) or sequencing (oligo 3). The sequences shown were synthesized on an Applied Biosystems, oligonucleotide 40 synthesizer and are complementary to the single stranded form of fd-tet (they are in the anti-sense form with respect to gene III). 4.2 shows the sequences of the various Constructs around the gene III insertion site. These sequences are drawn in the sense orientation with 45

respect to gene III; (A) fd-tet (and fdTSBst) (B) fdTPs/Bs and (C) fdTPs/Xh. The key restriction enzyme sites are shown along with the immunoglobulin amino acids contributed by the vectors, (amino acid single letter code is used, see Harlow, E., and Lane, D., 1988 supra.).

50 Figure 5 shows the nucleotide and amino acid sequences for scFv in the vector scFvD1.3 myc. This gives the sequence of the anti-lysoxyme single chain Fv

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and surrounding sequences in scPvD1.3 myc, showing the Nterminal pel B signal peptide sequence and the C-terminal myc tag sequence (Ward, E.S., et al., 1989, supra.). Also shown is the peptide sequence linking the VM and VL regions. The amino acid sequence is represented above the nucleotide sequence by the single letter code, see Harlow. E. and Lane D., 1988 supra.

Figure 6 shows the binding of pAbs to lysozyme and the effect of varying the amount of supernatant. Each point is the average of duplicate samples. Lysozyme was

10 point is the average of duplicate coated at 1 mg/ml in 50 mM NaHCO3.

Figure 7 shows the effect of varying the coating concentration of lysozyme or bovine serum albumin on the binding of pAbs to lysozyme in graphical form. Each

15 point is the average of duplicate samples.

Figure 8 shows the sequence around the cloning site in gene III of fd-CAT2. Restriction enzyme sites are shown as well as the amino acids encoded by antibody derived sequences. These are flanked at the 5' end by the gene III signal peptide and at the 3' end by alanine residues (encoded by the Not 1 restriction site) and the remainder of the mature gene III protein. The arrow shows the cleavage site for cutting of the signal peptide.

25 Figure 9 shows the binding of pAb (1.3) to lysczymes. Binding of phage as detected by ELISA to (a) hen egg-white lysczyme (HEL) (b) turkey egg-white lysczyme (TEL), (c) human lysczyme (HUL), (d) bovine serum albumin (BSA). A further control of (e) fdTPs/Bs 30 to HEL.

Figure 10 shows a map of FabD1.3 in pUC19.

Figure 11 shows the ELISA results providing a comparison of lysozyme-binding by phage-Fab and phage-scfv. Vector=fdCAT2 (example 5); fdscfv(OX)=pAbNQ11 (Example 9); fdVHCH1 (D1.3)=grown in normal cells (i.e. no L chain, see example 7); fdFab(D1.3) i.e. fdVHCH1 (D1.3) grown in cells containing D1.3 L chain; fdscfv

(D1.3)=pAbD1.3.

Figure 12 shows oligonuclectide probing of affinity purified phage. 10¹² phage in the ratio of 1 pAp (D1.3) in 4 x 10⁴ fdTPF/Bs phages were affinity purified and probed with an oligonuclectide specific for pAb (D1.3) A is a filter after one round of affinity purification (900 colondes total) and B is a filter after two rounds (372 colonies total).

Figure 13 shows the sequence of the anti-oxazolone antibody fragment NQ11 scFv. The sequence contributed by the linker is shown in the lower case. The sequence for VH is before the linker sequence and the sequence for VL

50 is after the linker.

Figure 14 shows the ELISA results for binding pAb NQ11 and pAb D1.3 and vector fdTFs/xh to specified

antigens.

Figure 15 shows the sequence surrounding the phoA insertion in fd-phoAlal66. The restriction sites used for cloning are shown, as well as the amino acids encoded by phoA around the insertion site. The first five amino

acids of the mature fusion come from gene III.

Figure 16(1) shows the structure of gene III and the native BamHI site into which a scFv coding sequence was inserted in example 13 and figure 16(2) shows the natural 10 peptide linker sites A and B for possible insertion of scrv coding sequences.

Figure 17 shows schematically the protocol for PCR assembly of mouse VH and VLK repertoires for phage

display described in example 14. 15

Figure 18 shows examples of the final products obtained with the procedure of example 14. Lanes a and b show the products of the initial PCR using heavy and light chain primers respectively; lane c shows the complete assembled 700bp product before final digestion with Not1 and ApaL1; M1, M2 markers \$174 Hae III digest and 123 base pair ladder (ERL Limited, P.O. Box 35,

Washington Road, Paisley, Scotland) respectively.
Figure 19 shows the binding of 1251-PDGF-BB to fd h-PDGFB-R phage in immunoprecipitation assay and comparison to fdTPs/Bs and no phage controls; binding is expressed as a percentage of the total 125 I-PDGF-BB added to the 25

incubation.

Figure 20 shows the displacement of 1251-ppgr-mg bound to fd-h-PDGFB-R phage using unlabelled PDGF-BB measured using an immunoprecipitation assay. Binding is expressed as a percentage of the total $^{125}\mathrm{r-ppGF-BB}$ added 30 to the incubation.

Figure 21 shows the displacement of 1251-ppgg-BB bound to fd-h-PDGFB-R phage using unlabelled PDGF-BB measured using an immunoprecipitation assay. Non-specific binding of 1251-PDGF-BB to vector phage fdTPs/Bs 35 in the absence of added unlabelled PDGF was deducted from each point.

Figure 22 shows the results of an ELISA of lysozyme 40 binding by pCAT-3 scFv D1.3 phagemid in comparison with pCAT-3 vector (both rescued by M13K07) and fdCAT2 scFv D1.3 as described in example 17. The ELISA was performed as described in example 6 with modifications detailed in example 18.

45 Figure 23 shows the digestion pattern seen when individual clones, selected at random from a library of single chain Fv antibody genes derived from an immunised mouse; are digested with BstNl.

Figure 24 shows VH and VK gene sequences derived 50 from the combinatorial library in example 21 and the hierarchical library in example 22,

Pigure 25 shows a matrix of ELISA signals for clones

derived from random combinatorial library. Designation of the clones is as in figure 24. The number of clones found with each combination is shown by the numerals.

Figure 26 shows a) the phagemid pHEN1 a derivative of pUC119 described in example 24; and b) the cloning

sites in the phagemid pHEN.

Figure 27. The antibody constructs cloned into fd-CAT2 and pHEN1 for display on the surface of phage. Constructs I, II, III and IV were cloned into both fd-CAT2 (as Apall-Not1 fragments) and pHEN1 (as Sfil-Not1

10 CAT2 (as ApalI-NotI fragments) and pHEN1 (as SfiI-NotI fragments) and pHEN1 (as SfiI-NotI fragments). All the constructs contained the heavy chain (VH) and light chain (VK) variable regions of the mouse anti-phOx antibody NQ10.12.5. The constant domains were human CK and CH1 (1) isotype).

Figure 28. Three ways of displaying antibody fragments on the surface of phage by fusion to gene III

protein.

Figure 29. Western blot of supernatant taken from pHEN1-II(+) or pHEN1(-) cultures in E.coli HB2151, showing secretion of Fab fragment from pHEN1-II only. The anti-human Fab detects both H and L chain. Due to the attached c-myc tag, the L chain, highlighted by both anti-c-myc tag and anti-human CK antisers, is slightly larger (calculated Mr 24625) than the H chain (c

Mr23145).

Figure 30 is a plot showing the effect of lysozyme dilution on ratio of ELISA signals obtained using pAbDl.3

or soluble scFv Dl.3.

or soluble sorv bits.

Figure 31 is a plot showing the effect of lysozyme dilution on ELISA signals obtained using fdTscFvDl.3 and soluble scFvDl.3.

ELISA screen of phage displaying soft fragments derived from the cell line 013 which express a monoclonal

35 from the cell line 013 which antibody directed against cestriol.

Figure 33 is a plot showing positive results from an ELISA screen of phage displaying scFv fragments derived from the cell line 014 which express a monoclonal

40 antibody directed against oestriol.

Figure 34 is a Western Blot showing expression of the alkaline phosphatase-gene 3 fusion. 16µl of 50 fold concentrate of each phage sample was detected on western blots with either anti-gene 3 antiserum (e-f) or with anti-alkaline phosphatase antiserum (c-f)

a) fd-phoAlal66 grown in TG1 cells

- b) fd-phoAla166 grown in KS272 cells
- c) fdCCAT2 grown in TG1 cells
- d) fdCAT2 grown in TG1 cells, mixed with 13 ng of

50 purified alkaline phosphatase

- e) fd-phoAlal66 grown in TG1 cells
- f) fdCAT2 grown in TG1 cells.

Figure 35 is a Western Blot showing ultrafiltration of phage-enzyme 100µl of 50 fold concentrate of phage (representing 5mls of culture supernatant) was centrifuged through ultrafiltration membranes with nominal molecular weight retention of 300,000 daltons. Western blots of flow through and retentate fractions were detected with anti-alkaline phosphatase antiserum. The equivalent of 800µl of original culture supernatant was run on the gel.

10 A. Phage were grown in TG1 cells. A. Phage were grown in TG1 cells. a) fd-phoAla166 before ultrafiltration (short exposure). b) fd-phoAla166 before ultrafiltration. c) fd-phoAlal66 material retained on ultrafiltration membrane.

Phage were grown in KS272 cells. a) fd-phoAlal66 before ultrafiltration. b) fd-phoAlal66 material 15 retained on ultrafiltration membrane. c) fdCAT2. d) fdCAT2 mixed with purified alkaline phosphatase before ultrafiltration. e) Retentate from sample d. f) Flow through from sample d.

20 Figure 36 Electrophoresis of samples from stages of a Fab assembly. Samples from different stages in the PCR Fab assembly process described in example 33 were subjected to electrophoresis on a 1% TAE-agarose gel.

Samples from a comparable scFv assembly process (as in 25 example 14) are shown for comparison. Samples left to

	right are:		
	M	300	Markers
	VHCHI	306	sequences encoding VHCH1 domains
30	VKCK		amplified by PCR
	AKCK	30 0	sequences encoding VKCK domains amplified by PCR
	~L	30.	Fab assembly reaction performed in absence of linkar
	. *		
35	+L	25	Fab PCR assembly reaction
			product VHCH1 plus VKCK plus
	M	ans.	Markers
	VΚ	*	sequences encoding VK domain amplified by PCR
40	VL		sequences encoding VH domains
			amplified by PCR
	-L	202	scrv assembly reaction in
	1.4		absence of linker
	+L	200	scFv assembly reaction in
45			presence of linker
	M	**	Markers

Figure 37. Comparison of ELISA signals with scFv D1.3 cloned in fd-CAT2 (fd) or pCAT-3. pCAT-3 scFv1.3 has 50 been rescued with M13K07 (KO7). M13K07AgIII No 3 (gIII No 3) or M13K07 gIIIANo 2 (g111No2). Phage antibodies are compared at 10 times (10x) 1 times (1x) or 0.1 times (0.1x) concentrations relative to concentration in the supernatant after overnight growth. The fdCAT2 and pCAT-3 non-recombinant vector signals were <0.01 at 10x concentration. M13K07 gIIIANo 1 did not rescue at all, as judged by no signal above background in this ELISA.

Figure 38. Western blot of PEG precipitated phage used in ELISA probed with anti-g3p. Free g3p and the c3p-scrp01.3 fusion bands are arrowed.

Sample 1 - fd scFvD1.3

10 Sample 2 - pCAT3 vector

Sample 3 - pCAT3 scFvDl.3 rescued with M13KO7, no IFTG

Sample 4 - pCAT3 scFvDl.3 rescued with M13K07, 50µM IPTG Sample 5 - pCAT3 scFvDl.3 rescued with M13K07, 100µM IPTG

Sample 6 - pCAT3 scFvDl.3 rescued with M13K07 gIII \(\Delta \) No3 (no IPTG)

Sample 7 - pCAT3 scfvDl.3 rescued with M13K07 gIII△ No 2 (no IPTG)

Panel A samples contain the equivalent of 8µl of phagemid culture supernatant per track, and 80µl of the 20 fd supernatant (10-fold lower phage yield than the phagemid). Panel B phagemid samples are those used in panel A at a five-fold higher sample loading (equivalent to 40µl of culture supernatant per track) to enable visualisation of the fusion band in samples rescued with parental M13KO7.

Figure 39 is a graph showing fdCAT2scFvDl.3 enrichment produced from a mixture of fdCAT2scFvDl.3 and

fdCAT2TPB4 by one round of panning.

Figure 40 is a graph showing fdCAT2scFvD1.3 on enrichment produced from a mixture of fdCAT2scFvD1.3 and fdCAT2TPB1 by one round of panning.

Figure 41. Western blot of phage proteins of fdCAT2(1) and fd-tet-SNase(2) with anti-g3p antiserum.

Marker molecular weights bands are indicated(kD).

Figure 42. Nuclease assay of soluble SNase (3 ng)(A-1),fd-tet-SNase(4 x 10 TU,(B-1),fd-CAT2(2 x 10 10 TU)(C-1) and of a PEG-precipitated fdCAT2 and SNase mixture(2 x 10 10 TU and 0.7ug)(D-1) in a 10-fold dilution series (1 to 3 or 4). Marker (M) is a HindIII digest of \-DNA(New 40 England Biolabs).

Figure 43. ELISA signals obtained with fd-tet, fd-CD4-V1 and fd-CD4-V1V2. In each group of three, the samples are left to right phage concentrate(SN); phage concentrate plus soluble CD4(SN + sCD4); phage

45 concentrate plus gp 120 (SN + gp 120).

Figure 44. shows the DNA sequence of scFv B18 (anti-

Figure 44. shows the DNA sequence of SCFV 518 (Enti

Figure 45 shows a map of the insert of sequences encoding FvD1.3 present in fd-tet FvD1.3 (example 39).

To designates the ribosome binding site. Gene III is now shown in its full length.

Figure 46. shows an ELISA assay of phages displaying

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FvD1.3 or scFvD1.3 by binding to plates coated with lysogyme. Signals obtained at various dilution factors are shown. FvD1.3 (AS-Stuffer) which does not express Fv was used as a control.

Figure 47. shows a schematic representation of steps involved in the PCR assembly of nucleotide sequences encoding human Fab fragments. Details are in example 40.

Figure 48. shows A. a map of plasmid pJM1-Fabbl.3 . which is used for the expression of soluble human Fab 10 fragments and as a template for the synthesis of linker DNA for Fab assembly. B. a schematic representation of sequences encoding a Fab construct. C. The sequence of DNA template for the synthesis of linker DNA for Fab assembly. 15

Figure 49. shows a schmatic representation of steps involved in the PCR assembly of nucleotide sequences encoding human scFv fragments. Details are in example 42.

Figure 50. ELISA assay of phage antibodies using 20 plates coated with turkey egg lysogyme. Two clones Bi and A4 are shown derived by mutagenesis and selection from pAbD1.3 (example 45). Concentration (x axis) refers to the concentration of phage for each sample relative to the concentration in culture supernatant. El has raised 25 binding to turkey egg lysogyme compared to D1.3. A4 has

reduced binding to hen egg lysogyme compared to D1.3. Figure 51. ELISA of phage antibodies binding to HEL and TEL. Clone 1 is fdCAT2scFvD1.3. Clones 2 to 10 were obtained from the library (example 46) after selection.

The background values as defined by binding of these clones to BSA were subtracted.

Figure 52. shows the DNA sequence of the light chains D1.3 M1F and M21 derived by selection from a hierarchical library in example 46.

Figure 53 shows a Fv lambda expression vector (example 48) derived from pUC119. It contains the 35 rearranged lambdal germ line gene. The heavy and light chain cassettes each contain a ribosome binding site upstream of the pel B leader (Restriction sites shown as: 40 H=Hind III; Sp=SphI; B=BamHI, E=EcoRI,

Materials and Methods

The following procedures used by the present applicants are described in Sambrook, J. et al., 1989 supra.: restriction digestion, ligation, preparation of competent cells (Hanahan method), transformation. analysis of restriction enzyme digestion products on agarose gels, purification of DNA using phenol/chloroform, 5'-end labelling of oligonucleorides, filter screening of bacterial colonies, preparation of 2xTY medium and plates, preparation of tetracycline stock

solutions, PAGE of proteins, preparation of phosphate buffered saline.

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All enzymes were supplied by New England Biolabs (CP Laboratories, PO Box 22, Bishop's Stortford, Herts., England) and were used according to manufacturer's instructions unless otherwise stated.

The vector fd-tet (Zacher, A.N. et al., 1980, supra) was obtained from the American Type Culture Collection (ATCC No. 37000) and transformed into competent TG1 cells (genotype: K125 (lac-pro), sup E, thi, hsdD5/F traD36, pro A+B+, Lac 19, lac 6M15).

Viral particles were prepared by growing TG1 cells 10 containing the desired construct in 10 to 100 mls 2xTY medium with 15 µg/ml tetracycline for 16-24 hours. culture supernatant was collected by centrifugation for 10 mins at 10,000 rpm in an 8 x 50 ml rotor, Sorval RC-5B

centrifuge. Phage particles were precipitated by adding 1/5th volume 20% polyethylene glycol (PEG)/2.5M NaCl and These were spun for 15 leaving at 4°C for 1 hour. minutes as described above and the pellets resuspended in 10 mM Tris/HCl pH 8, 1mm EDTA to 1/100th of the original volume. Residual bacteria and undissolved material were 20 removed by spinning for 2 minutes in a microcentrifuge. Single stranded DNA for mutagenesis or sequencing was prepared from concentrated phage according to Sambrook,

J., et al., 1989,, supra.

Index of Examples Example I Design of Insertion Point Linkers and Construction of Vectors

This example covers the construction of two derivatives of the phage vector fd-tet: a) fdTPs/Bs for the insertion of VH coding sequences; and b) fdTPs/Xh for 30 the insertion of scrv coding sequences. The derivative vectors have a new BstEII site for insertion of sequences.

Example 2 Insertion of Immunoglobulin Fv Domain into Phage 35

This example covers the insertion of scFv coding sequences derived from an anti-lysozyme antibody D1.3 into fdTPs/Xh to give the construct fdTscFvD1.3. Example 3 Insertion of Immunoglobulin VH Domain into

40 Phage This example covers the insertion of VH coding sequences derived from an anti-lysozyme antibody D1.3 into fdTPs/Bs to give the construct fdTVHD1.3. Example 4 Analysis of Binding Specificity of Phage

Antibodies This example investigates the binding specificities

of the constructs fdTscFvDl.3 and fdTVHDl.3. Example 5 Construction of fdCAT2

This example covers the construction of the derivative fdCAT2 of the phage vector fdTPs/Xh. 50 derivative has restriction sites for enzymes that cut DNA infrequently.

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Example 6 Specific Binding of Phage Antibody (pAb) to Antigen

This example shows the binding of pAb fdTscFvD1.3 to lysozyme by ELISA.

Example 7 Expression of FabD1.3

This example concerns the display of an antibody Fab fragment at the phage surface. The VH-CH1 chain is The VL-CL chain is expressed by expressed by fdCAT2. DUC19 in a bacterial host cell also infected with fdCAT2.

10 Example 8 Isolation of Specific, Desired Phage from a Mixture of Vector Phage

This example shows how a phage (e.g. fdTscFvD1.3) displaying a binding molecule can be isolated from vector phage by affinity techniques.

15 Example 9 Construction of pAb Expressing Anti-Hapten Activity

This example concerns the insertion of scFv coding sequences derived from the anti-oxazolone antibody NQ11 into fdTPs/Xh to generate the construct pAbNQ11. 20 example shows the binding of pAbNQ11 to oxazalone by ELISA.

Example 10 Enrichment of pAbD1.3 from Mixtures of other

pabs by Affinity Purification This example shows how a phage (eg. pAbD1.3)

displaying one sort of biding molecule can be isolated 25 from phage (e.g. pAbNQll) displaying another sort of binding molecule by affinity techniques. Example 11 Insertion of a Gene Encoding an Enzyme

(Alkaline Phosphate) into fdCAT2

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This example concerns the invention of coding sequences for an enzyme into the vector fdCAT2 to give the phage enzyme, fdphoAlall6, Example 12 Measuring Enzyme Activity Phage - Enzyme

This example shows the functionality of an enzyme (alkaline phosphatase) when displayed at the phage surface (fdphoAla166).

Example 13 Insertion of Binding Molecules into Alternative Sites in the Phage

This example covers the insertion of scrv coding 40 sequences derived from a) the anti-lysozyme antibody D1.3; and b) the anti-oxazalone antibody NO11 into a BamH1 site of fdTPs/Xh to give the constructs fdTBam1 having an NO11 insert.

Example 14 PCR Assembly of Mouse VH and VLK Repertoires 45

for Phage Display

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This example concerns a system for the display on phage of all VH and VLK repertoires encoded by a mouse. The system involves the following steps. 1) Preparation of RNA from spleen. 2) Preparation of cDNA from the RNA 3) Use of primers specific for antibody sequences to PCF amplify all VH and VLK cDNA coding sequences 4) Use of PCR to create a linker molecule from linking pairs of VH

and VLK sequences 5) Use of PCR to assemble continuous DNA molecules each comprising a VH sequence, a linker and a VLK sequence. The specific VH/VLK combination is randomly derived 6) Use of PCR to introduce restriction

5 Example 15. Insertion of the Extracellular Domain of a Human Receptor for Platelet Derived Growth Factor (PDGF)

Isoform BB into fdCAT2

This example concerns the insertion of coding sequences for the extracellular domain of the human receptor for 10 PDGF into the vector fdCAT2 to give the construct fdhPDGFBR.

Example 16. Binding of 125 I-PDGF-BB to the Extracellular Domain of the Human Receptor for PDGF Isoform BB Displayed on the Surface of fd Phage. Measured using an Displayed on the Surface of fd Phage.

15 Immunopredipitation Assay.

This example shows that the human receptor PDGF Isoform BB is displayed on the surface of the phage in a form which has the ability to bind its ligand.

Example 17. Construction of Phagemid Containing Gene III 20 Fused with the Coding Sequence for a Binding Molecule. This example concerns the construction of two phagemids based on pUC119 which separately contain gene III from

fdCAT2 and the gene III scFv fusion fdCAT2seFvDI.3 to generate pCAT2 and pCAT3 scFvD1.3 respectively. 25

Example 18. Rescue of Anti-Lysozyme Antibody Specificity

from pCAT3scFvD1.3 by M13KO7 This example describes the rescue of the coding sequence for the gene IIIscFv fusion from pCAT3scFvD1.3 by M13M07 helper phage growth, phage were shown to be displaying 30 scrv anti-lypozyme activity by ELISA.

Example 19. Transformation Efficiency of FCAT-3 and pCAT-3 scFvDl.3 Phagemids

This example compared the efficiency of the phagemids pVC119, pCAT-3 and pCAT3scFvD1.3 and the phage fdCAT2 to 35 transform E.coli.

Example 20 PCR Assembly of a Single Chain Fv Library from

an Immunised Mouse

This example concerns a system for the display on phage of scFv (comprising VH and VL) from an immunised mouse 40 using the basic technique outlined in example 14 (cDNA preparation and PCR assembly of the mouse VH and VLK repertoires) and ligating the PCR assembled sequences into fdCAT2 to create a phage library of 105 clones.

Testing of 500 clones showed that none showed specificity 45

against phox.

Selection of Antibodies Specific for 2-Example 21. phenyl-5-oxazolone from a Repertoire from an Immunised

This example shows that phage grown from the library 50 established in example 20 can be subjected to affinity selection using phOX to select those phage displaying

scFv with the desired specificity.

Example 22. Generation of Further Antibody Specificities by the Assembly of Hierarchial Libraries.

This example concerns the construction of hierarchial libraries in which a given VM sequence is combined with the complete VLK repertoire and a given VLK sequence is

combined with the complete VH repertoire and selection from these libraries of novel VH and VL pairings.

Example 23. Selection of Antibodies Displayed on

10 Bacteriophage with Different Affinities for 2-phenyl-5oxazolone using Affinity Chromatography

This example concerns the separation by affinity techniques of phages displaying scFv fragments with differing binding affinities for a given antiquen.

Example 24. Construction of Phagemid pHEN1 for the Expression of Antibody Fragments Expressed on the surface of Bacteriophage following Superinfection

This example concerns the construction of the phagemid pHEN1 derived from pUC119. pHEN1 has the features shown in Fig. 26.

Example 25. Display of Single Chain Fv and Fab Fragments
Derived from the Anti-Oxazolone Antibody NO 10.12.5 on
Bacteriophage fd using pHEN1 and fdCAT2.

This example describes the display of scFv and Fab fragment with a specificity against phOx on the surface of a bacteriophage. For display of scFv the phagemidd pHEN1 comprises the sequences encoding scFv (VH and VL) for rescue by either the phages VSM13 or fdCAT2. For display of Fab the phage fdCAT2 comprises the sequence for either the H or L chain as a fusion with g3p and the

30 for either the H or L chain as a fusion with g3p and the phagemid pHEN1 comprises the sequence for the appropriate H or L chain partner.

Example 26. Rescue of Phagemid Encoding a Gene III protein Fusion with Antibody Heavy or Light Chains by Phage Encoding the Complementary Antibody Displayed on Phage and the Use of this Technique to make Dual Combinatorial Libraries

This example covers the use of phage antibodies encoding the antibody heavy or light chain to rescue a phagemid encoding a gene 3 protein fusion with the complementary chain and the assay of Fab fragments displayed on phage in ELISA. The use of this technique in the preparation of a dual combinatorial library is discussed.

45 Example 27 Induction of Soluble scFv and Fab Fragments using Phagemid pHEN1

This example covers the generation of soluble scFv and Fab fragments from gene III fusions with sequences encoding these fragments by expression of clones in pHEN1 in an E.coli strain which does not suppress amber mutations.

Example 28 Increased Sensitivity in ELISA of Lysozyme

using fdTscFvDl.3 as Primary Antibody compared to Scluble scFvD1.3

This example covers the use of fdTscFvD1.3 in ELISA showing that lower amounts of lysozyme can be detected with phage antibody fdTscFvDi.3 than with soluble scFvD1.3.

Example 29 Direct Rescue and Expression of Mouse Monoclonal Antibodies as Single Chain Fv Fragments on the

Surface of Bacteriophage fd

This example covers the display on phage as 10 functional scFv fragments of two clones directly derived from cells expressing monoclonal antibodies directed Both clones were established to be against oestriol. functional using ELISA.

Example 30 Kinetic Properties of Alkaline Phosphatase 15

Displayed on the Surface of Bacteriophage fd

This example concerns the demonstration that the kinetic properties of an enzyme, alkaline phosphatase, displayed on phage are qualitatively similar to those of

the same enzyme when in solution. 20

Example 31 Demonstration using Ultrafiltration that Cloned Alkaline Phosphatase Behaves as Part of the Virus Particle

This example concerns the construction of the phage enzyme fdphoArgl66 and the demonstration that both the 25 fusion protein made and the catalytic activity observed derive from the phage particle.

Example 32 Affinity Chromatography of Phage Alkaline

Phosphatase

This example concerns the binding of alkaline 30 phosphatase displayed on phage to an arsenate-Sepharose affinity column and specific elution of these phage using the reaction product, phosphate.

Example 33 PCR Assembly of DNA Encoding the Fab Fragment of an Antibody Directed against Oxazolone

This example covers the construction of a DNA insert encoding a Fab fragment by separate amplification of heavy and light chain DNA sequences followed by assembly. The construct was then inserted into the phage vector fdCAT2 and the phagemid vector pHEN1 and the Fab fragment

displayed on the surface was shown to be functional. Example 34 Construction of a Gene III Deficient Helper

Phage

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This example describes the construction of a helper phage derived from M13KO7 by deleting sequences in gene 45 Rescue of pCAT3-scFvDl.3 is described. scFvD1.3 is expressed at a high level as a fusion using the deletion phage, equivalent to expression using fdCAT2-scFvD1.3.

Example 35 Selection of bacteriophage expressing scFv 50 fragments directed against lysozyme from mixtures

according to affinity using a panning procedure

This example concerns the selection of bacteriophage according to the affinity of the sofv fragment directed against lysozyme which is expressed on their surface. The phage of different affinities were bound to Petri

- dishes coated with lyszzyme and, following washing, bound phage eluted using triethylamine. Conditions were found where substantial enrichment could be obtained for a phage with a 5-fold higher affinity than the phage with which it was mixed.
- 10 Example 36 Expression of Catalytically Active Staphylococcal Nuclease on the Surface of Bacteriophage fd
- This example concerns the construction of a phage enzyme which expresses Staphylococcal nuclease and the demonstration that the phage enzyme retains nuclease activity.
 - Example 37 Display of the Two Aminoterminal Domains of Human CD4 on the Surface of fd Phage
- This example covers the cloning of genes for domains of CD4, a cell surface receptor and member of the immunoglobulin superfamily, into bacteriophage fd. The receptor is shown to be functional on the surface of phage by binding to the HIV protein gp120.
- Example 38 Generation and Selection of Mutants of an Anti-4-hydroxy-3-nitrophenylacetic acid (NP) Antibody expressed on Phage using Mutantor strains
- This example covers the introduction of mutations into a gene for an antibody cloned in phage by growth of the phage in strains which randomly mutate DNA due to defects in DNA replication. Several mutations are introduced into phage which can then be selected from
- parent phage.

 Example 39 Expression of a Fy Fragment on the Surface of Bacteriophage by Non-Covalent Association of VH and VL domains
 - This example shows that functional Fv fragments can be expressed on the surface of bacteriophage by non-covalent association of VH and VL domains. The VH domain is expressed as a gene III fusion and the VL domain as a soluble polypeptide. Sequences allowing expression of these domains from the anti-lysozyme antibody Dl.3 in this form were introduced into phage and the resulting displayed Fv fragment shown to be functional by ELISA.
- Example 40 A PCR Based Technique for one step Cloning of Human V-genes as Fab Constructs
- This example gives methods for the assembly of Fab fragments from genes for antibodies. Examples are given for genes for antibodies directed against Rhesus-D in a human hybridoma and a polyclonal lymphoblestic cell line. Example 41 Selection of Phage Displaying a Human Fab Fragment directed against the Rhesus-D Antigen by binding to Cells displaying the Rhesus D Antigen on their Surface

This example concerns the construction of, and display of phage antibodies from, a phagemid encoding a human Fab fragment directed against the Rhesus D antigen. Phage displaying this antigen were then affinity selected from a background of phage displaying scFvDL.3 antilysozyme on the basis of binding to Rhesus-D positive red blood cells.

Example 42 A PCR Based Technique for One Step Cloning of

Human scFv Constructs

This example describes the generation of libraries of sofv fragments derived from an unimmunized human.

Examples are given of the preparation for phage display of libraries in phagemids of sofv fragments derived from LuG and IoM sequences.

15 Example 43 Isolation of Binding Activities from a Library

of scFvs from an Unimmunized Human

This example describes the isolation, from the library of scfv fragments derived from IgM genes of an unimmunized human, of clones for phage antibodies directed against BSA, lysozyme and oxazolone. Selection was by panning or affinity chromatography and analysis of binding specificity by ELISA. Sequencing of the clones showed them to be of human origin.

Example 44 Rescue of human IgM library using helper phage

25 Tacking gene 3 (g3)

This example covers the isolation, from the library of scFv fragments of unimmunized human IgM genes, of clones of phage antibodies of clones for phage antibodies specific for thyroglobulin and oxazolome. In this example rescue was with MISKO7gIII No3 (NCTC12478), a

example rescue was with habygill has their rounds of selection appeared necessary for a phagemid library rescued with this phage compared to one rescued with MISKOT.

35 Example 45 Alteration of Fine Specificity of scFvDl.3 displayed on Phage by Mutagenesis and Selection on

Immobilized Turkey Lysozyme

This example covers the in vitro mutagenesis of pCATscPvDl.3 by replacement, with random amino acids, of residues known to be of importance in the preferential recognition of hen egg lysozyme over turkey egg lysozyme by scFvDl.3. Following selection for phage antibodies recognising turkey egg lysozyme by affinity chromatography, clones were analyzed for specificity by ELSA. Two groups of clones were found with more equal

45 ELISA. Two groups of clones were found with more equal recognition of hen and turkey lysozymes, one with increased ELISA signal with the turkey enzyme and one with reduced signal for the hen enzyme.

Example 46 Modification of the Specificity of an Antibody by Replacement of the VLK Domain by a VLK Library derived

from an Unimmunised Mouse

This example shows that replacement of the VL domain

of scFvD1.3 specific for hen eggwhite lysozyme (HEL) with a library of VL domains allows selection of scFv fragments which bind also to turkey eggwhite lysozyme The scFv fragments were displayed on phage and selection by panning on tubes coated with TEL. Analysis by ELISA showed clones with enhanced binding to TEL compared to HEL. Those with highest binding to TEL were

sequenced. Example 47 Selection of a Phage Antibody Specificity by binding to an Antigen attached to Magnetic Beads. Use of 10 a Cleavable Reagent to allow Elution of Bound Phage under Mild Conditions

This examples covers the use of a cleavable bond in the affinity selection method to allw release of bound 15 phage under mild conditions. pAbNQ11 was enriched approximately 600 fold from a mixture with pAbD1.3 by selection using biotinylated Ox-BSA bound to magnetic beads. The cleavage of a bond between BSA and the biotin allows elution of the phage.

Example 48 Use of Cell Selection to provide an Enriched 20 Pool of Antigen Specific Antibody Genes, Application to reducing the Complexity of Repertoires of Antibody Fragments Displayed on the Surface of Bacteriophage

This example covers the use of cell selection to produce an enriched pool of genes encoding antibodies 25

directed against 4-hydroxy-3-nitrophenylacetic acid and describes how this technique could be used to reduce the complexity of antibody repertoires displayed on the surface of bacteriophage. 30 Example 1

Design of Insertion Point Linkers and Construction of Vectors

The vector fd-tet has two BstEII restriction sites flanking the tetracycline resistance gene (fig 3). Since 35 the strategy for inserting the VH fragments was to ligate them into a newly inserted BstEII site within gene III, it was advantageous to delete the original BstEII sites from fd-tet. This was achieved by digesting fd-tet with the restriction enzyme BstEII, filling-in the 5'

40 overhangs and re-ligating to generate the vector fdToBst. Digestion of fd-tet with BstEII (0.5 units/µl) was carried out in 1x KGB buffer (100 mM potassium glutamate. 23 mM Tris-acetate (pH 7.5), 10 mM magnesium acetate, 50

µg/ml bovine serum albumin, 0.5 mM dithiothreitol (Sambrook, J., et al., 1989, supra.) with DNA at a 45 concentration of 25 ng/ul. The 5' overhang was filled in, using 2x KGB buffer, 250 µM each dNTP's (Pharmacia

Ltd., Pharmacia House, Midsummer Boulevard, Milton Keynes, Bucks., UK.) and Klenow Fragment (Amersham International, Lincoln Place, Green End, Aylesbury, Bucks., UK) at 0.04 units/pl. After incubating for 1 50 hour at room temperature, DNA was extracted with

phenol/chloroform and precipitated with ethanol.

Ligations were carried out at a DNA concentration of 50ng/µl). Ligations were transformed into competent TG1 cells and plated onto TY plates supplemented with 15 µg/ml tetracycline. This selects for vectors where the gene for tetracycline resistance protein has reinserted into the vector during the ligation step. Colonies were picked into 25 mls of 2xTY medium supplemented with 15 µg/ml tetracycline and grown overnight at 37°C.

Double stranded DNA was purified form the resulting clones using the gene-clean II kit (Biol01 Inc., PO Box 2284, La Jolla, California, 92038-2284, USA.) and according to the small scale rapid plasmid DNA isolation procedure described therein. The orientation of 5 of the resulting clones was checked using the restriction enzyme Clal. A clone was chosen which gave the same pattern of restriction by Clal as fd-tet, but which had no BstE II

sites.

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In vitro mutagenesis of fdToBst was used to generate vectors having appropriate restriction sites that facilitate cloning of antibody fragments downstream of the gene III signal peptide and in frame with the gene III coding sequence. The oligonucleotide directed mutagenesis system version 2 (Amersham International) was used with oligo 1 (figure 4) to create fdTPs/Bs (to facilitate cloning of VH fragments). The sequence offdTPs/Bs (figure 4) was confirmed using the sequence version 2.0 kit (USB Corp., PO Box 22400, Cleveland, Ohio, 44122, USA.) with oligo 3 (figure 4) as a primer.

30 A second vector fdTPs/Xh (to facilitate cloning of single chain Fv fragments) was generated by mutaganising fdTPs/Bs with oligo 2 according to the method of Venkitaraman, A.R., Nucl. Acid Res. 17, p 3314. The sequence of fdTPs/Xh (figure 4) was confirmed using the sequence version 2.0 kit (USB Corp.) with oligo 3 as a

primer.

Clearly, alternative constructions will be apparent to those skilled in the art. For example, M13 and/or its host becteria could be modified such that its gene III could be disrupted without the onset of excessive cell death; the modified fd gene III, or other modified protein, could be incorporated into a plasmid containing a single stranded phage replication origin, such as pUC119, superinfection with modified phage such as KO7 would then result in the encapsulation of the phage antibody genome in a coat partially derived from the helper phage and partly from the phage antibody gene III construct.

The detailed construction of a vector such as fdTPs/Bs is only one way of achieving the end of a phage antibody. For example, techniques such as sticky feet cloning/mutagenesis (Clackson, T. and Winter, G. 1989

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Nucl. Acids. Res., $\underline{17}$, p 10163-10170) could be used to avoid use of restriction enzyme digests and/or ligation steps. Example 2.

5 Insertion of Immunoglobulin Fv Domain into Phage

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The plasmid scFv Dl.3 myc (gift from g. Winter and A. Griffiths) contains VH and VL sequences from the antibody Dl.3 fused via a peptide linker sequence to form a single chain Fv version of antibody Dl.3. The sequence of the scFv and surrounding sequences in scFvDl.3 myc is shown in figure 5.

The $D\bar{1}.3$ antibody is directed against hen egg lysozyme (Harper, M. et al., 1987, Molec. Immunol. 24, 97-108) and the scFv form expressed in E.coli has the same specificity (A. Griffiths and G. Winter personal

Communication).

Digestion of scFv D1.3 myc with Pstl and Xhol (these restriction sites are shown on Fig. 5), excises a fragment of 693 bp which encodes the bulk of the scFv. Ligation of this fragment into fdTFe/Xh cleaved with Pstl and Xhol gave rise to the construct fdTscFvD1.3 encoding the gene III signal peptide and first amino acid fused to the complete D1.3 scFv, followed by the mature gene III protein from amino acid 2.

25 The vector fdTPs/Xh was prepared for ligation by digesting with the Pstl and Xhol for 2 hours followed by digestion with calf intestinal alkaline phosphatase (Boehringer Mannheim UK Ltd., Bell Lane, Lewes, East Sussex, BN7 lLG) at one unit/ul for 30 minutes at 37°C. Fresh calf intestinal alkaline phosphatase was added to a final total concentration of 2 units/ul and incubated for a further 30 minutes at 37°C. The reaction was extracted

ethanol and dissolved in water. The insert from scFVD1.3

myc was excised with the appropriate restriction enzymes
(PstI and XhoI) extracted twice with phenol/chloroform,
precipitated with ethanol and dissolved in water.
Ligations were carried out as described in example 1,
except both vector and insert samples were at a final
concentration of 5 ng/ul each. The formation of the
correct construct was confirmed by sequencing as

three times with phenol/chloroform, precipitated with

described in example 1.

To demonstrate that proteins of the expected size were produced, virions were concentrated by PEG precipitation as described above. The samples were prepared for electrophoresis as described in Sambrook J. et al 1989 supra. The equivalent of Zmls of supernatant was loaded onto an 18% SDS polyacrylamide gel. After electrophoresis, the gel was soaked in gel running buffer (50 mM tris, 380 mM Glycine, 0.1%SDS) with 20% methanol

for 15 minutes. Transfer to nitrocellulose filter was executed in fresh lx running buffer/20% methanol using

TE70 Semi Phor a semi-dry blotting apparatus (Hoeffer, 654 Minnesota Street, Box 77387, San Francisco, California 94107, USA.).

AFter transfer, the filter was blocked by incubation for 1 hour in a 2% solution of milk powder (Marvel) in phosphate buffered saline (PBS). Detection of scFv and VH protein sequences in the phage antibody fusion proteins was effected by soaking the filter for 1 hour with a 1/1000 dilution (in 2% milk powder) of a rabbit polyclonal antiserum raised against affinity purified, 10 bacterially expressed scFv fragment (gift from G. Winter). After washing with PBS (3 x 5 minute washes), bound primary antibody was detected using an anti-rabbit 15

antibody conjugated to horseradish peroxidase (Sigma, Fancy Road, Poole, Dorset, BH17 7NH, UK.) for 1 hour. The filter was washed in PES/0.1% triton X-100 and developed with 0.5 mg/ml 3,3'-diaminobenzidine tetrahydrochloride (DAB), 0.02% cobalt chloride, 0.03%

hydrogen peroxide in PBS.

The results showed that with clones fdTVHD1.3 (from 20 example 3 incorporating sequences coding for VH) and fdTscFvD1.3 (incorporating sequences coding for scFv) a protein of between 69,000 and 92,500 daltons is detected by the anti-Fv serum. This is the expected size for the fusion proteins constructed. This product is not 25 observed in supernatants derived from fd-tet, fdT&Bst or

fdTPs/Xh. Example 3.

Insertion of Immunoglobulin VH Domain into Phage Antibody The VH fragment from D1.3 was generated from the 30 plasmid pSW1-VHD1.3-TAG1 (Ward, E.S. et al., 1989 supra.). Digestion of this plasmid with Pstl and BstEII

generates the fragment shown between positions 113 and 432 in figure 5. Cloning of this fragment into the Pstl and BatEII sites of fdTPs/Bs gave rise to the construct fdTVHD1.3 which encodes a fusion protein with a complete VH domain inserted between the first and third amino acids of the mature gene III protein (amino acid two has

been deleted).

The methods used were exactly as in example 2 except 40 that the vector used was fdTPs/Bs digested with Pstl and BstEII.

Example 4.

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Analysis of Binding Specificity of Phage Antibodies

The binding of the various phage antibodies to the specific antigen, lysozyme, was analysed using ELISA Phage antibodies (e.g. fdTVHD1.3 and techniques. fdTsc/FvD1.3) were grown in E.coli and Phage antibody particles were precipitated with PEG as described in the materials and methods. Bound phage antibody particles were detected using polyclonal sheep serum raised against the closely related phage M13.

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ELISA plates were prepared by coating 96 well plates (Falcon Microtest III flexible plate. Falcon: Becton Dickinson Labware, 1950 Williams Drive, Oxnard. California, 93030, USA.) with 200 ul of a solution of lysozyme (1mg/ml unless otherwise stated) in 50 mm NaHCO3 for 16-24 hours. Before use, this solution was removed, the plate rinsed several times in PBS and incubated with 200 ul of 2% milk powder/PBS for 1 hour. After rinsing several times with PBS, 100 ul of the test samples were added and incubated for 1 hour. Plates were washed (3 10 rinses in 0.05% Tween 20/PBS followed by 3 rinses in PBS alone). Bound phage antibodies were detected by adding 200 ul/well of a 1/1000 dilution of sheep anti-M13 polyclonal antiserum (gift from G. Winter, although an 1.5 equivalent antibody can be readily made by one skilled in the art using standard methodologies) in 2% milk powder/PBS and incubating for 1 hour. After washing as above, plates were incubated with biotinylated anti-sheep antibody (Amersham International) for 30 minutes. Plates 20 were washed as above, and incubated with streptavidinhorseradish peroxidase complex (Amersham International). After a final wash as above. 0.5 mg/ml ABTS substrate in citrate buffer was added (ABTS = 2'2'-azinobis (3ethylbenzthiazoline sulphonic acid); citrate buffer * 50 25 mM citric acid, 50 mM tri-sodium citrate at a ratio of 54:46. Hydrogen peroxide was added to a final concentration of 0.003% and the plates incubated for 1 The optical density at 405 nm was read in a Titertek multiskan plate reader.

Figure 6 shows the effect of varying the amount of phage antibody. 100 ul of various dilutions of PEG precipitated phage were applied and the amount expressed in terms of the original culture volume from which it was derived. Signals derived from both the scFv containing phage antibody (fdTscFvD1.3) and the VH containing phage antibody (fdTVHD1.3) and the VH containing phage antibody were higher than that derived from the phage antibody vector (fdTPs/Xh). The highest signal to noise ratio occurs using the equivalent of 1.3 mls of culture.

Figure 7 shows the results of coating the plates with varying concentrations of lysozyme or bovine serum albumin (BSA). The equivalent of 1 ml of the original phage antibody culture supernatant was used. The signals from supernatants derived from fdTscFvDl.3 were again higher than those derived from fdTPs/Xh when lysozyme coated wells were used. There was no significant difference between these two types of supernatant when the plates were coated with BSA. Broadly speaking the level of signal on the plates is proportional to the amount of lysozyme coated. These results demonstrate that the binding detected is specific for lysozyme as the

antigen.

Example 5:

Construction of fd CAT 2

It would be useful to design vectors that enable the use of restriction enzymes that cut DNA infrequently, thus avoiding unwanted digestion of the antibody gene inserts within their coding sequence. Enzymes with an eight base recognition sequence are particularly useful in this respect, for example Notl and Sfil. Chaudhary et al (PNAS 87 p1066-1070, 1990) have identified a number of restriction sites which occur rarely in antibody variable genes. The applicant has designed and constructed a vector that utilises two of these sites, as an example of how this type of enzyme can be used. Essentially sites

for the enzymes Apall and Notl were engineered into fdTFs/Xh to create fdCAT2.

The oligonucleotide: 5'ACT TTC AAC AGT TTC TGC GGC CGC CGC TTT GAT CTC GAG CTC CTG CAG TTG GAC CTG TGC ACT GTG AGA ATA GAA 3'

was synthesised (supra fig 4 legend) and used to mutagenise fdTPs/Xh using an in vitro mutagenesis kit from Amersham International as described in example 1, to create fd-CAT2. The sequence of fd-CAT2 was checked around the site of manipulation by DNA sequencing. The final sequence around the insertion point within gene III

25 is shown in figure 8. N.B. fdCAT2 is also referred to herein by the alternative

terminologies fd-tet-DOG1 and fdDOG1.

Example 6
Specific Binding of Phage-antibody (pAb) to Antigen

The binding of pab D1.3 (fdTscFvD1.3 of example 2) to lysozyme was further analysed by ELISA.

Methods.
1. Phage growth.

Cultures of phage transduced bacteria were prepared in 10-100 mls 2 x TY medium with 15 pg/ml tetracycline and grown with shaking at 37°C for 16-24 hrs. Phage supernatant was prepared by centrifugation of the culture (10 min at 10,000 rpm, 8 x 50 ml rotor, Sorval RC-58 centrifuge). At this stage, the phage titre was 1 - 5 x 1010/ml transducing units. The phage were precipitated by adding 1/5 volume 20% PEG 2.5 M NaCl, leaving for 1 hr

by adding 1/5 volume 20% PEG 2.5 M NaCl, leaving for 1 hr at 4°C, and centrifuging (supra). The phage pellets were resuspended in 10 mM Tris-Hcl, 1mM EDTA PH 8.0 to 1/100th of the original volume, and residual bacteria and aggregated phage removed by centrifugation for 2 min in a

bench microcentrifuge. ELISA

Plates were coated with antigen (1 mg/ml antigen) and blocked as described in example 4. 2 x 10₁₀ phage transducing units were added to the antigen coated plates in phosphate buffered saline (PBS) containing 2* skimmed milk powder (MPBS). Plates were washed between each step

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with three rinses of 0.5% Tween-20 in PBS followed by three rinses of PBS. Bound phage was developed by incubating with sheep anti-M13 antisers and detected with horseradish peroxidase (HRP) conjugated anti-goat serum (Sigma, Poole, Dorset, UK) which also detects sheep immunoglobulins and ABTS (2'2'-azinobis (3-ethylbenzthiazoline sulphonic acid). Readings were taken at 405 nm after a suitable period. The results (figure 9) show that the antibody bearing-phage had the same pattern of reactivity as the original D1.3 antibody (Harper, M., Lema, F., Boulot, G., and Poljak, F.J. (1987) Molec. Immunol. 24, 97-108), and bound to hen egg-white lysozyme, but not to turkey egg-white lysozyme, human lysozyme or bovine serum albumin. The specificity of the phage is particularly illustrated by the lack of binding to the turkey egg-white lysozyme that differs

from hen egg-white lysozyme by only 7 amino acids.

Example 7.

Expression of Fab D1.3 The aim of this example was to demonstrate that the scFv format used in example 2 was only one way of displaying antibody fragments in the pAb system. A more commonly used antibody fragment is the Fab fragment-(figure 1) and this example describes the construction of a pab that expresses a Fab-like fragment on its surface and shows that it binds specifically to its antigen. The applicant chose to express the heavy chain of the antibody fragment consisting of the VH and CH1 domains from coding sequences within the pAb itself and to coexpress the light chain in the bacterial host cell infected with the pAb. The VH and CHI regions of antilysozyme antibody D1.3 were cloned in fd CAT2, and the corresponding light chain cloned in plasmid pUC19. The work of Skerra and Pluckthun (Science 240, p1038-1040 (1988) and Better et al 1988 supra; demonstrated that multimeric antigen binding fragments of the antibody molecule could be secreted into the periplasm of the bacterial cell in a functional form using suitable signal sequences. However, in these publications, special measures were described as being needed to recover the binding protein from the cell, for example Skerra and Pluckham needed to recover the Fv fragment from the periplasm by affinity chromatography. The present applicants have shown that it is possible to direct the binding molecule to the outside of the cell on a phage particle, a process that requires several events to occur: correct secretion and folding of the binding molecule; association of the chains of the binding molecule; correct assembly of the phage particle; and export of the intact phage particle from the cell.

Alternatively, it is possible however, to express the light chain from within the pAb genome by, for ×

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example, cloning an expression cassette into a suitable place in the phage genome. Such a suitable place would be the intergenic region which houses the multicloning sites engineered into derivative of the related phage M13 (see, for example, Yanisch-Perron, C. et al., Gene 33,

p103-119, (1985)). The starting point for this example was the clone Fab D1.3 in pUC19, a map of which is shown in figure 10. The regions hybridising with the oligonucleotides KSJ6 and 7 below are shown underlined in fig 10. The sequence encoding the VH-CH1 region (defined at the 5' and 3' edges by the oligonucleotides KSJ6 and 7 below) was PCR amplified from Fab Dl.3 in pUC19 using oligonucleotides KSJ 6 and 7, which retain the Pst I site at the 5' end and introduce a Xho I site at the 3' end, to facilitate cloning into fd CAT2. The sequences for the

oligonuclectides KSJ6 and 7 are shown below. underlined region of KSJ7 shows the portion hybridising with the sequence for D1.3.

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KSJ6:5' AGG TGC AGC TGC AGG AGT CAG G 3' KSJ7: 5' GGT GAC CTC GAG TGA AGA TTT GGG CTC AAC TTT C 3' PCR conditions were as described in example II, except that thirty cycles of PCR amplification were performed with denaturation at 92°C for 45 seconds, annealing at 55°C for 1 minute and extension at 72°C for 1 minute. The template used was DNA from TG1 cells containing Fab D1.3 in pUC19 resuspended in water and boiled. The template DNA was prepared from the colonies by picking some colony material into 100µl of distilled H₂O and boiling for 10 mins. lul of this mixture was used in a 20ul PCR. This regime resulted in amplification of the expected fragment of approximately 600bp. This fragment was cut with Pst I and Xho I, purified from an agarose gel and ligated into Pst 1/Xho 1-cut fdCAT2. The PCR mixture was extracted with phenol/chloroform and ethanol precipitated (Sambrook et al. supra.) before digestion

35 with Pstl and Xhol (New England Biolabs according manufacturers recommendations. The fragment was resolved on 1% Tris-Acetate EDTA agarose gel (Sambrook et al. supra) and purified using Geneclean (BIO 101, Geneclean, 40 La Jolla, San Diego, California, USA) according to manufacturers recommendations.

fd-CAT2 vector DNA was digested with Pst 1 and Xho 1 (New England BioLabs) according to manufacturers recommendations, extracted with phenol/chloroform and ethanol precipitated (Sambrook et al. supra.).

75ng of Pst 1/Xho 1-digested vector DNA was ligated to 40ng of PCR-amplified Pstl /Xho I-digested hEGF-R fragment in 12µ1 of ligation buffer (66mM TrisHC1 (pH7.6), 5mM MgCl₂, 5mM dithiothreitol, (100µg/ml bovine serum albumin, 0.5mM ATP, 0.5mM Spermidine) and 400 units T4 DNA ligase (New England BioLabs) for 16 hours at 16°C.

Two µl of the ligation mixture was transformed into 200µl of competent E.coli MC1061 cells, plated on 2TY agar containing 15µg/ml tetracycline and incubated at 30°C for 20 hours. A portion of the ligation reaction mixture was transformed into E.coli MC1061 (Available from, for example Clontech Laboratories Inc, Palo Alto, California) and colonies identified by hybridisation with the oligonucleotide Dl.3CDR3A as described in example 10.

The presence of the VHCH1 gene fragment was likewise confirmed by PCR, using Oligonucleotides KSJ6 and 7. A representative clone was called fd CATZVHCH1 Dl.3. The heavy chain was deleted from Fab Dl.3 in pUC19 by Sph I cleavage of Fab Dl.3 plasmid DNA. The pUC 19 2.7Kb fragment containing the light chain gene was purified

15 from a TAE agarose gel, and long of this DNA self-ligated and transformed into competent E.coll TG1. Cells were plated on 2TY agar containing ampicillin (100µg/ml) and incubated at 30°C overnight. The resulting colonies were used to make miniprep DNA (Sambrook et al. supra), and the absence of the heavy chain gene confirmed by digestion with Sph I and Hind III. A representative

clone was called LCD1.3 DHC.

An overnight culture of fd CAT2VHCH1 D1.3 cells was microcentrifuged at 13,000Xg for 10 minutes and 50µl of the supernatant containing phage particles added to 50µl of an overnight culture of LCD1.3 DHC cells. The cells were incubated at 37°C for 10 minutes and plated on 2TY agar containing ampicillin (100µg/ml) and 15µg/ml tetracycline. Phage were prepared from some of the 30 resulting colonies and assayed for their ability to bind lysozyme as described in example.6.

The results (Figure 11) showed that when the heavy and light chain Fab derivatives from the original antibody D1.3 were present, the pAb bound to lysozyme.

35 pAb expressing the fd VHCHI fragment did not bind to lysozyme unless grown in cells also expressing the light chain. This shows that a functional Fab fragment was produced by an association of the free light chain with VHCHI fragment fused to gene III and expressed on the

40 surface of the pAb. Example 8

Isolation of Specific, Desired Phage from a Mixture of Vector Phage.

The applicant purified pAb (D1.3) (criginally called fdTscFVD1.3 in example 2) from mixtures using antigen affinity columns. pAb (D1.3) was mixed with vector fd phage (see table 1) and approximately 10¹² phage passed over a column of lysozyme-Sepharose (prepared from cyanogen bromide activated sepharose 4B (Pharmacia, Milton Keynes, Bucks, UK.) according to the manufacturers instructions. TG1 cells were infected with appropriate dilutions of the elutes and the colonies derived, were

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analysed by probing with an oligonucleotide that detects only the pAb (D1.3) see Table 1 and Fig. 12. A thousand fold enrichment of pab(D1.3) was seen with a single column pass. By growing the enriched phage and passing it down the column again, enrichments of up to a million fold were seen.

Enrichment was also demonstrated using purely immunological criteria. For example, 1012 phage (at a ratio of 1 pAb (D1.3) to 4 x 106 fdTPs/Bs) was subjected to two rounds of affinity selection, and then 25 colonies picked and grown overnight. The phage was then assayed for lysozyme binding by ELISA (as example 6). colonies yielded phage with lysozyme binding activities, see table 1, and these were shown to encode the scrv (D1.3) by PCR screening (example 13, using 30 cycles of 1 minute at 92°C, 1 minute at 60°C, 1 minute at 72°C using

CDR3PCR1 and oligo 3 (fig. 4) as primers). Thus very rare pabs can be fished out of large populations, by using antigen to select and then screen

the phage. 20

In this example, affinity chromatography of pAbs and oligonucleotide probing were carried out as described below. .

Approximately 1012 phage particles in 1ml MPBS were loaded onto a 1 ml lysozyme-Sepharose affinity column 25 which had been prewashed in MPBS. The column was washed in turn with 10 ml PBS; then 10 ml 50 mM Tris-HCl, 500 mM Nacl pH 7.5; then 10ml 50 mM Tris-HCl 500 mM NaCl pH 8.5; then 5 mls 50 mM Tris-HCl, 500 mM NaCl pH 9.5 (adjusted with triethylamine) and then eluted with 5 ml 100 mM 30 triethylamine. The eluate was neutralised with 0.5 M sodium phosphate buffer pH 6.8 and the phage plated for analysis. For a second round of affinity chromatography, the first column eluste was plated to about 30,000

After overnight growth, colonies per petri dish. 35 colonies were then scraped into 5 ml 2 x TY medium, and a 20 ul aliquot diluted into 10 ml fresh medium and grown overnight. The phage was PEG precipitated as described above, resuspended in 1 ml MPBS and loaded onto the

column, washed and eluted as above. 40

Oligonucleotides sythesised: CDR3PCR1 5'TGA GGA C(A or T) C(A or T) GC CGT CTA CTA CTG TGC 3'

40 pmole of oligonucleotide VHIFOR (Ward, E. S., et al (1989) Nature 341, 544-546), specific to pAb (D1.3) 45 was phosphorylated with 100 μC1 α-32P ATP, hybridised (lpmole/ml) to nitrocellulose filters at 67°C in 6 x saline sodium citrate (SSC) Sambrook et al., supra. buffer for 30 minutes and allowed to cool to room temperature for 30 mins, washed 3 x 1 min at 60°C in 0.1 50 x SSC.

Example 9

Construction of pAb Expressing Anti-hapten Activity

Oxazolone is a hapten that is commonly used for studying the details of the immune response. The anti-oxazalone antibody, NQ11 has been described previously (E. Gherardi, R. Pannell, C. Milstein, J. Immunol. Method 126 51-68). A plasmid containing the VM and VL gene of NQ11 was converted to a scFv form by inserting the BstEII/SacI fragment of scFvPll.3 myc (nucleotides 432-499 of Fig. 5) between the VH and VL genes to generate pscFvNQ11, the sequence of which is shown in fig. 13. This scFv was cloned into the PstI/Xhol site of FdTFs/Xh (as described

sequence of which is shown in fig. 13. This scFv was cloned into the Pstl/Xhol site of FdTPs/Xh (as described earlier) to generate pAb NQl1 has an internal Pstl site and so it was necessary to do a complete digest of pscFvNQl1 with Xhol followed by a partial digest with 15 Pstl).

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The specific binding of pAb NQ11 was confirmed using ELISA. ELISA plates were coated at 37°C in 50 mM NAHCO3 at a protein concentration of 200 µg/ml. Plates were coated with either hen egg lysozyme (HEL), bovine serum

20 albumin (BSA), or BSA conjugated to oxazolone (OX-BSA) (method of conjugation in Makela O., Kartinen M., Pelkonen J.L.T., Karjalainen K. (1978) J. Exp. Med.148 1644). Preparation of phage, binding to ELISA plates, washing and detection was as described in example 6.

25 Samples were assayed in duplicate and the average absorbance after 10 minutes presented in figure 14.

This result demonstrates that the pah NQ1 binds the correct antigen. Figure 14 also shows that pab D1.3 and pab NQ1 bind only to the antigen against which the

30 original antibodies were raised., Example 10

Enrichment of pAb D1.3 from Mixtures of Other pAb by Affinity Purification

3 x 10¹⁰ phage in 10 mls of PBSM at the ratios of pAb D1.3 to pAb NQ11 shown in table 2 were passed over a 1 ml lysozyme Sepharose column. Washing, elution and other methods were as described in example 8 unless otherwise stated. Eluates from the columns were used to infect TG1 cells which were then plated out. Colonies

40 were probed with a probe which distinguishes pAb D1.3 from pAb NQ11. The sequence of this oligonucleotide (D1.3CDR3A) is:-

5'GTA GTC AAG CCT ATA ATC TCT CTC 3'

Table 2 presents the data from this experiment. An enrichment of almost 1000 fold was achieved in one round and an enrichment of over a million fold in two rounds of

purification. This parallels the result described in example 8. $\tt Example \ 11$

50 Insertion of a Gene Encoding an Enzyme (Alkaline phosphatase) into fd-CAT2

As an example of the expression of a functional

enzyme on the bacteriophage surface, the applicants have chosen bacterial alkaline phosphatase, an enzyme that normally functions as a dimer (McCracken, S. and Meighen, E., J. Biol. Chem. 255, p2396-2404, (1980)). The oligonucleotides were designed to generate a PCR product with an Apa LI site at the 5' end of phoA gene and a Not I site at its 3' end, thus facilitating cloning into fd-CAT 2 to create a gene III fusion protein. The

oligonuclectides synthesised were: 10 phoAl:5' TAT TCT CAC AGT GCA CAA ACT GTT GAA CGG ACA CCA GAA ATG CCT GTT CTG 3' and,

phoA2:5' ACA TGT ACA TGC GGC CGC TTT CAG CCC CAG AGC GGC

TTT C3'
The sequence of the phoA gene is presented in Chang C. N.
et al., Gene 44, p121-125 (1986). The plasmid amplified
(pEK86) contains an alkaline phosphate gene which differs
from the sequence of Chang et al. by a mutation which

converts arginine to alamine at position 166.

The PCR reaction was carried out in 100µl of 10 mM
Tris/HCl pH 8.3, containing 50 mM KCl, 5mMGNTP 2.5 mM
MgCl₂, 0.01½ gelatin, 0.25 units/µl of Taq polymerase
(Cetus/Perkin Elmer) and 0.5µg/ml template. The template
was the pEK86 plasmid (described by Chaidaroglou et al.,
Biochemistry 27 p8338-8343, 1988). The PCR was carried
out in a Techne (Techne, Duxford, Cambridge, UK) PHC-2
dri-block using thirty cycles of 1 min at 92°C, 2 min at
50°C, 3 min at 72°C.

The resultant product was extracted with phenolichloroform, precipitated with ethanol, and the pellet dissolved in 35µl water. Digestion with 0.3 units/µl of Apa Ll was carried out in 150µl volume according to manufacturers instructions for two hours at 37°C. After heat inactivation of the enzyme at 65°C, NaCl was added to a final concentration of 150mM and 0.4

35 units/µl Noti enryme added. After incubation for 2 hours at 37°C, the dispest was extracted with phenolichloroform and precipitated as above, before being dissolved in 30µl of water. The vector fd-CAT2 was sequentially dispeted with Apa Ll and Noti according to the manufacturers instructions and treated with calf intestinal alkaline phosphatase as described in example 2. The sample was

extracted three times with phenol:chloroform, precipitated with ethanol and dissolved in water. The ligations were performed with a final DNA concentration of 1-2ng/µl of both the cut fd-CAT2 and the digested PCR product. The ligations were transformed into competent TG1 cells and plated on 2xTY tet plates. Identification

of clones containing the desired insert was by analytical PCR performed using the conditions and primers above, on boiled samples of the resulting colonies. The correct clone containing the phoA gene fused in frame to g.ne III was called fd-phoAla 166. The sequence at the junction

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of the cloning region is given in figure 15. Example 12

Measuring Enzyme Activity of Phage-enzyme

Overnight cultures of TG1 or KS272 (E.coli cells lacking phoA. Strauch K. L., and Beckwith J. PNAS 85 1576-1580, 1988) cells containing either fd-phoAla 166 or fd-CAT2 were grown at 37°C in 2xTY with 15ug/ml tetracycline. Concentrated, PEG precipitated phage were prepared as described earlier. Enzyme assays (Malamy, M.H. and Horecker B.L., Biochemistry 3, p1893-1897, 3.0

(1964)) were carried out at 24°C in a final concentration of 1M Tris/HC1 pH 8.0, 1mm 4-nitrophenyl phosphate (Sigma), 1mm MgC12. 100µl of a two times concentrate of this reaction mixture was mixed with 100ul of the test sample in a 96 well plate. Absorbance readings were taken every minute for 30 minutes at a wavelength of 405nm in a Titretek Mk 2 plate reader. Initial reaction rates were calculated from the rate of change of

absorbance using a molar absorbance of 17000 1/mol/cm. Standard curves (amount of enzyme vs. rate of change of absorbance) were prepared using dilutions of purified bacterial alkaline phosphatase (Sigma type III) in 10mM Tris/HC1 pH 8.0, 1mM EDTA. The number of enzyme Tris/HC1 pH 8.0, 1mM EDTA. The number of enzyme molecules in the phage samples were estimated from the actual rates of change of absorbance of the phage samples

25 and comparison to this standard curve.

The results in Table 3 show that alkaline phosphatase activity was detected in PEG precipitated material in the sample containing fd-phoAlal66 but not 30 fd-CAT2. Furthermore, the level of activity was consistent with the expected number of 1-2 dimer molecules of enzyme per phage. The level of enzyme activity detected was not dependent on the host used for growth. In particular, fd-phoAla166 grown on phoA minus 35 hosts showed alkaline phosphatase activity.

Therefore, the phage expressed active alkaline phosphatase enzyme, from the phoA-gene III fusion, on the phage surface.

Example 13

40 Insertion of Binding Molecules into Alternative Sites in the Phage

The availability of an alternative site in the phage for the insertion of binding molecules would open up the possibility of more easily expressing more than one binding molecule e.g. an antibody fragment in a single DAb. This may be used to generate single or multiple binding specificities. The presence of two distinct binding activities on a single molecule will greatly increase the utility and specificity of this molecule. It may be useful in the binding of viruses with a high mutational rate such as human immunodeficiency virus. In

addition, it may be used to bring antigens into close

proximity (e.g. drug targetting or cell fusion) or it may act as a "molecular clamp" in chemical, immunological

or enzymatic processes.

The vector fd-tet and the derivatives described here, have a single BamHl site in gene 3. This has previously been used for the expression of peptide fragments on the surface of filamentous bacteriophage (Smith GP. (1985) Science 228 p1315-1317 and de la Cruz et al. (1988) J Biol. Chem. 263 p4318-4322). provides a potential alternative site for the insertion 10

of antibody fragments.

DNA fragments encoding scFv's from Dl.3 or NQ11 were generated by PCR using the primers shown below. primers were designed to generate a fragment with BamHl sites near both the terminii, to enable cloning into the 15 BamHl site of gene3 (see figure 16(1)). cligonucleotides used, also ensure that the resulting PCR product lacks Pstl and Xhol restriction sites normally used for manipulating the scFv's (see figure 16(1)). This will facilitate subsequent manipulation of a second 20

antibody fragment in the usual way at the N terminus of gene 3. The oligonucleotides used were:-G3Baml 5'TTT AAT GAG GAT CCA CAG GTG CAG CTG CAA GAG 3'

G3Bam2 5'AAC GAA TGG ATC CCG TTT GAT CTC AAG CTT 3'.

Preparation of vector and PCR insert 25 The FCR reaction was carried out in an 80 µl reaction as described in example 11 using lng/µl of template and 0.25U/µl of Tag polymerase and a cycle regime of 94°C for 1 minute, 60°C for 1 minute and 70°C for 2 minutes over 30 cycles. The template was either 30 pscFvNQll (example 9) or scFvDl.3 myc (example 2). Reaction products were extracted with phenol:chloroform, precipitated, dissolved in water and digested with BamH1 according to manufacturers instructions. The digest was re-extracted with phenol: chloroform, precipitated and 35

dissolved in water. The vector fdTPs/Xh was cleaved with BamHl and treated with calf intestinal phosphatase and purified as described in example 2. Ligations were set up at a vector concentration of approximately 6ng/µl and a PCR 40 insert concentration of approximately 3ng/µl. These were ligated for 2.5 hours at room temperature before transforming into competent TG1 cells and plating on TY The resultant colonies were probed as tet plates.

described in example 8. DNA was prepared from a number 45 of colonies and the correct orientation and insert size confirmed by restriction digestion with Hind III in isolation or in combination with BamHl. (One Hind III site is contributed by one of the primers and the other 50 by the vector).

Two clones containing a D1.3 insert (fdTBaml) and fdTBam2) and one containing an NQ11 insert (NQ11Bam1)

were grown up and phage prepared as described earlier. ELISAs were carried out as described in example 6. No specific signal was found for any of these clones suggesting that the natural BamHl site is not a suitable site for insertion of a functional antibody (results not shown).

It may be possible to clone into alternative sites to retain binding activity. The peptide repeats present in gene III may provide such a site (figure 16 blocks A and B). This can be done by inserting a BamHl site and using the PCR product described above. To facilitate this, the natural BamHl site was removed by mutagenesis with the oligonucleotide GSmut6Bam shown below (using an in vitro mutagenesis kit (Amersham International)):-

15 G3mut6Bam 5' CA AAC GAA TGG GTC CTC CTC ATT A 3'
The underlined residue replaces an A residue, thereby removing the BamHl site. DNA was prepared from a number of clones and several mutants lacking BamHl sites

identified by restriction digestion.

The oligonucleotide G3 Bamlink was designed to introduce a BamHl site at a number of possible sites within the peptide linker sites A and B, see figure 16(2). The sequence of the linker is: Bamlink 5°CC (G or A) CC ACC CTC GGA TCC (G or A) CC ACC

25 CTC 3'

Its relationship to the peptide repeats in gene III is shown in figure 16.

Example 14 PCR Assembly of Mouse VH and VL Kappa (VLK) Repertoires

30 for Phage Display

The principle is illustrated in figure 17. Details are provided in sections A to F below but the broad outline is first discussed.

cDNA is prepared from spleen RNA from an appropriate 35 mouse and the VH and VLK repertories individually amplified. Separately, primers reverse and complementary to VHIFOR-2 (domain 1) and VLKZBACK (domain 2) are used to amplify an existing scFvcontaining DNA by PCR. (The term FOR refers to e.g. 40 a primer for amplification of sequences on the sense strand resulting in antisense coding sequences. The term BACK refers to e.g. a primer for amplification of sequences on the antisense strand resulting in sense coding sequences). This generates a 'linker' 45 molecule encoding the linker with the amino acid sequence (1 letter code) (GGGGS)3 which overlaps the two primary (VH and VLK) PCR products.

2. The separate amplified VH, Vik and linker sequences now have to be assembled into a continuous DNA molecule by use of an 'assembly' PCR. In the secondary 'assembly' PCR, the VH, VLK and linker bands are combined and assembled by virtue of the

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above referred to overlaps. This generates an assembled DNA fragment that will direct the expression of VH and one VLK domain. The specific VH/VLK combination is derived randomly from the separate VH and VLK repertoires referred to above.

The assembly PCR is carried out in two stages. Firstly, 7 rounds of cycling with just the three bands present in the PCR, followed by a further 20 rounds in the presence of the flanking primers VHIBACK (referring to domain 1 of VH) and VLKFOR. The nucleotide sequences for these oligonucleotide primers are provided under the section entitled 'Primer Sequences' below. This two stage process, avoids the potential problem of preferential amplification of the first combinations to

15 be assembled. For cloning into the phage system, the assembled

repertoires must be 'tagged' with the appropriate restriction sites. In the example provided below this is illustrated by providing an ApaLl restriction site at the 20 VH end of the continuous DNA molecule and a Not 1 site at the VLK end of the molecule. This is carried out by a third stage PCR using tagged primers. The nucleotide sequences for these oligonucleotide primers are also provided under the section entitled 'Primer Sequences' below. There are however, 4 possible kappa light chain sequences (whereas a single consensus heavy chain sequence can be used). Therefore 4 oligonucleotide primer sequences are provided for VLK.

For this third stage PCR, sets of primers which create the new restriction site and have a further 10 nucleotides on the 5' side of the restriction site have been used. However, long tags may give better cutting, in which case 15-20 nucleotide overhangs could be used.

Scrupulously clean procedures must be used at all times to avoid contemination during PCR. Negative controls containing no DNA must always be included to monitor for contamination. Gel boxes must be depurinated. A dedicated Geneclean kit (810 101, Geneclean, La Jolla, San Diego, California, USA) can be used according to manufacturers instructions to extract DNA from an agarose gel. The beads, NaI and the NEW Wesh

should be aliquoted.
All enzymes were obtained from CP Laboratories, P.O.
Box 22, Bishop's Stortford, Herts CM20 3DH and the
manufacturers recommended and supplied buffers were used
unless otherwise stated.

A. RNA Preparation

RNA can be prepared using may procedures well known to those skilled in the art. As an example, the following protocol (Triton X-100 lysis, phenol/SDS RNase inactivation) gives excellent results with spleen and hybridoma cells (the addition of VRC (veronal ribosyl

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complex) as an RNase inhibitor is necessary for spleen cells). Guanidinium isothiocyanate/CsCl procedures (yielding total cellular RNA) also give good results but are more time-consuming.

- 5 1. Harvest 1 to 5 x 107 cells by centrifugation in a bench tope centrifuge at 800xg for 10 minutes at 4°C. Resuspend gently in 50ml of cold PBS buffer. Centrifuge the cells again at 800xg for 10 minutes at 4°C, and discard supernatant.
- 10 2. On ice, add 1 ml ice-cold lysis buffer to the pellet and resuspend it with a lml Gilson pepette by gently pepetting up and down. Leave on ice for 5 minutes.

 After lysis, remove cell debris by centrifuging at 1300 rpm for 5 minutes in a microfuge at 4°C, in

15 precooled tubes. 4. Transfer 0.5 ml

4. Transfer 0.5 ml of the supernatant to each of two eppendorfs containing 50µl 10% (w/v) SDS and 250 µl phenol (previously equilibrated with 100 mM Tris-HCl pH 8.0). Vortex hard for 2 minutes, then microfuge (13000 rpm) for five minutes at room temperature. Transfer the upper supernature.

Transfer the upper, aqueous, phase to a fresh tube.

5. Re-extract the aqueous upper phase five times with

- 0.5 ml of phenol.

 6. Precipitate with 1/10 volume 3M sodium acetate and
 25 2.5 volumes ethanol at 20°C overnight or dry iceisopropanol for 30 minutes.
 - 7. Wash the RNA pellet and resuspended in 50 µl to check concentration by 00260 and check 2 µg on a 1% agarose gel. 40µg of RNA was obtained from spleen cells derived from mice.

Lysis buffer is [10mM Tris-HC1 pH 7.4, lmM MgC12, 150mM NaCl, 10mM VRC (New England Biolabs), 0.5% (w/v) Triton X-100], prepared fresh.

35 Lysis buffer is [10mM Tris-HCl pH 7.4, lmM MgCl₂, 150mM NaCl, 10mM VRC (New England Biolabs), 0.5% (w/v) Triton X-1001, prepared fresh.

B. cDNA Preparation

cDNA can be prepared using many procedures well to known to those skilled in the art. As an example, the following protocol can be used:

1. Set up the following reverse transcription mix:

DEPC is diethylpyrocarbonate, the function of which is to inactivate any enzymes that could decrade DNA

or RNA

ii) dNTP is deoxynucleotide triphosphate

iii) DTT is dithiothreitol the function of which is as an antioxidant to create the reducing environment necessary for enzyme function.

iv) RNasin is a ribonuclease inhibitor obtained from Promega Corporation, 2800 Woods Hollow Road,

Madison, Wisconsin, USA.

2. Dilute 10 µg RNA to 40 µl final volume with DEPCtreated water. Heat at 65°C for 3 minutes and hold on ice for one minute (to remove secondary structure).

3. Add to the RNA the reverse transcription mix (58 µl) and 4 µl of the cloned reverse transcriptase 'Super RT' (Anglian Biotech Ltd., Whitehall House, Whitehall Road, Colchester, Essex) and incubate at 42°C for one hour.

 Boil the reaction mix for three minutes, cool on ice for one minute and then spin in a microfuge to pellet debris. Transfer the supernatant to a new tube.

10 x first strand buffer is [1.4M KCl, 0.5M Tris-HCl

pH 8.1 at 42°C 80mM MgCl2].

The primers anneal to the 3' end. Examples of kappa Light chain primers are MUKIFONX, MJKZFONX, MJKZFONX and MJKSFONX (provided under 'Primer Sequences' below) and examples of heavy chain primers are MIGGI, 2 (CTG GAC AGG GAT CCA GAG TTC CA) and MIGG3 (CTG GAC AGG GCT CCA TAG TTC CA) which anneal to CH1.

30 Alternatively, any primer that binds to the 3' end of the variable regions VH, VLK, VL, or to the constant

regions CH1, CK or CL can be used. C. Primary PCRs

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For each PCR and negative control, the following reactions are set up (e.g. one reaction for each of the four VLKs and four VH PCRs). In the following, the Vent DNA polymerase sold by (C.P. Laboratories Ltd (New England Biolabs) address given above) was used. The buffers are as provided by C.P. Laboratories.

H ₂ O 10 x Vent	huffer	32.5 5
20 x Vent		2.5
5mM dNTPs		1.5
FOR prime:	r 10 pmol/µl)	2.5
	er 10pmol/ul	2.5

The FOR and BACK primers are given in the section below entitled 'Primer Sequences'. For VH, the FOR primer is VHIFOR-2 and the BACK primer is VHIBACK. For VLK the FOR primers are MJKIFONX, MJKZFONX, MJKAFOKX and MJKSFONX (for the four respective kappa light chains) and the BACK

primer is VKZBACK. Only one kappa light chain BACK primer is necessary, because binding is to a nucleotide sequence common to the four kappa light chains.

UV this mix 5 minutes. Add 2.5 ul cDNA preparation (from E above), 2 drops paraffin oil (Sigma Chemicals, Poole, Dorset, UK). Place on a cycling heating block, e.g. PHC-2 manufactured by Techne Ltd. Duxford UE, preset at 94°C. Add lul Vent DNA polymerase under the paraffin. Amplify using 25 cycles of 94°C 1 min, 72°C 2

10 min. Post-treat at 60°C for 5 min.

Purify on a 2% lmp (low melting point agarose/TAE (tris-acctate EDTA)gel and extract the DNA to 20 µl H₂0 per original PCR using a Geneclean kit (see earlier; in accordance with the manufacturers instructions.

15 D. Preparation of linker Sat up in bulk (e.g. 10 times)

	H ₂ O	34.3
	10 x Vent buffer	5
20	20 x Vent BSA	2.5
	5mM dNTPs	2
	LINKFOR primer 10 pmol/ul)	2.5
	LINKBACK primer 10pmol/ul	2.5
	DNA from fcFv D1.3 (example 2)	1
25	Vent enzyme	0.2

The FOR and BACK primers are given in the section below entitled 'Primer Sequences'. The FOR primer is LINKFOR and the BACK primer is LINKFORCK. Cover with paraffin and place on the cycling heating block (see above) at 94°C. Amplify using 25 cycles of 94°C 1 min, 65°C 1 min, 72°C 2

min. Post-treat at 60°C for 5 min.

Purify on 2% imp/TAE gel (using a loading dye without bromophenol blue as a 93bp fragment is desired) and elute with SPIN-X column (Costar Limited, 205 Broadway, Cambridge, Ma. USA.,) and precipitation. Take up in 5 µl H₂O per PCR reaction. E. Assembly PCRs

A quarter of each PCR reaction product (Sul) is 40 used for each assembly. The total volume is 25ul.

For each of the four VLK primers, the following are set up:

H₂O 4.95 45 10 x Vent buffer 2.5 20 x Vent BSA 2.25 5mM dNTPs 0.8

UV irradiate this mix for 8 min. Add 5ul each of Vh and VK band from the primary PCRs and 1.5 ul of linker as isolated from the preparative gels and extracted using the Geneclean kit as described in C and D above. Cover

with paraffin. Place on the cycling heating block preset at 94°C . Add hil Vent under the paraffin. Amplify using 7 cycles of 94°C 2 min, 72°C 4 min. Then return the temperature to 94°C .

Add 1.5ul each of VH1BACK and the appropriate VKFOF primers MJK1FONX, MJKZFONX, MJKZFONX or MJK5FONX (10 pmol/µl) at 94°C. The primers should have been UV-treated as above. Amplify using 20 cycles of 94°C 1.5 min, 72°C 2.5 min. Post-treat at 66°C for 5 min. Purify on 2% 1mp/TAE gel and extract the DNA to 20µl H₂O per assembly PCR using a Geneclean kit (see earlier) in

accordance with the manufacturers instructions. F. Adding Restriction Sites

For each assembly and control set up:

15		nī.
	H ₂ O	36.5
	10 x Tag buffer	5
	5mM dNTPs	2
	FOR primer (10 pmol/µl)	2.5
20	BACK primer (10 pmol/pl)	2.5
	Assembly product	1

The FOR and BACK primers are given in the section below entitled 'Primer Sequences'. The FOR primer is any of JKINOTIO, JKZNOTIO, JKZNOTIO or JKSNOTIO (for the four respective kappa light chains) for putting a Notl restriction site at the VLK end. The BACK primer is HEKAPAIO for putting an Apall restriction site at the VH end.

So Cover with paraffin and place on the cycling heating block preset at 94°C. Add 0.5 µl Cetus Taq DNA polymerase (Cetus/perkin-Elmer, Beaconsfield, Bucks, UK) under the paraffin. Amplification is carried out using 11 to 15 rounds of cycling (depends on efficiency) at 94°C l min, 55°C l min, 72°C 2 min. Post-treat at 60°C for 5 min.

10 x Tag buffer is [0.1M Tris-HCl pH 8.3 at 25°C, 0.5M KCl. 15mM MgCl., lmg/ml gelatin].

G. Work-up

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40 Purify once with CHCl3/IAA (isoamylalcohol), once with phenol, once with CHCl3/IAA and back-extract everything to ensure minimal losses. Precipitate and wash twice in 70% EtOH. Dissolve in 70ul H₂O.

The DNA (joined sequence) above refers to the assembled 50 DNA sequence comprising in the 5' to 3' direction

Apail restriction site

VA sequence

Linker sequence VLK sequence

Not 1 restriction site.

The VLK sequence may be any one of four possible kappa chain sequences.

The enzymes Not 1 above. Apall below and the buffers NEB Not 1. NEB BSA above and the NEB buffer 4 (below) are obtainable from CP Laboratories. New England Biolabs mentioned above.

10 Re-precipitate, take up in 80µl H20. Add to this 10ul NEB buffer 4 and 10ul Apal 1.

Add the enzyme Apall in aliquots throughout the day, as it has a short half-life at 37°C.

Purify on 2% lmp/TAE gel and extract the DNA using a 15 Geneclean kit, in accordance with the manufacturers instructions. Redigest if desired.

Final DNA product The final DNA product is an approximate 700 bp fragment with Apa Ll and Notl compatible ends consisting of randomly associated heavy and light chain sequences linked by a linker. A typical molecule of this type is the scFvD1.3 molecule incorporated into fdscFvD1.3 described in example 3. These molecules can then be ligated into suitable fd derived vectors, e.g. fdCAT2

(example 5), using standard techniques. 25

Primer sequences
Primery PCR oligos (restrictions sites underlined): VHIFOR-2 TGA GGA GAC GGT GAC CGT GGT CCC TTG GCC CC VHIBACK AGG TSM ARC TGC AGS AGT CWG G

MJKIFONX CCG TTT GAT TTC CAG CTT GGT GCC 30 MJKZFONX CCG TTT TAT TTC CAG CTT GGT CCC
MJKZFONX CCG TTT TAT TTC CAA CTT TGT CCC
MJKZFONX CCG TTT CAG CTC CAG CTT GGT CCC

VKZBACK GAC ATT GAG CTC ACC CAG TCT CCA 35 Ambiguity codes M = A or C, R = A or G, S = G or C. W = A or T

PCR oligos to make linker: LINKFOR TGG AGA CTC GGT GAG CTC AAT GTC LINKBACK GGG ACC ACG GTC ACC GTC TCC TCA

For adding restriction sites: HEKAPA10 CAT GAC CAC AGT GCA CAG GTS MAR CTG CAG SAG TOW GG

JKINOT10 GAG TCA TTC TGC GGC CGC CCG TTT GAT TTC CAG CTT 45 GGT GCC JK2NOT10 GAG TCA TTC TGC GGC CGC CCG TTT TAT TTC CAG CTT GGT CCC

JK4NOT10 GAG TCA TTC TGC GGC CGC CCG ITT TAT ITC CAA CTT TGT CCC

50 JK5NOT10 GAG TOA THE TGE GGE CGE CCG THE CAG CTE CAG CTE GGT CCC

Example 15

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Insertion of the Extracellular Domain of a Human Receptor for Platelet Derived Growth Factor (PDGF) soform BB into fd CAT2

A gene fragment encoding the extracellular domain of the human receptor for platelet derived growth factor isoform BB (h-PDGFB-R) was isolated by amplification, using the polymerase chain reaction, of plasmid RP41, (from the American Type Culture collection, Cat. No.50735), a cDNA clone encoding amino-acids 43 to 925 of the PDGF-B receptor (Gronwald, R.G.K. et al PNAS 85 10 p3435-3439 (1988)). Amino acids 1 to 32 of h-PDGFB-R constitute the signal peptide. The oligonucleotide primers were designed to amplify the region of the h-PDGFB-R gene corresponding to amino acids 43 to 531 of the encoded protein. The primer RPDGF3 for the N-terminal region also included bases encoding amino acids 15 33 to 42 of the h-PDGFB-R protein (corresponding to the first ten amino acids from the N-terminus of the mature

protein) to enable expression of the complete extracellular domain. The primers also incorporate a 20 unique Apall site at the N-terminal end of the fragment and a unique Xhol site at the C terminal end to facilitate cloning into the vector fdCAT2. The sequence

of the primers is:

RPDGF3 5' CAC AGT GCA CTG GTC GTC ACA CCC CCG GGG CCA GAG CTT GTC CTC AAT GTC TCC AGC ACC TTC GTT CTG 3' RPDGFZ 5' GAT CTC GAG CTT AAA GGG CAA GGA GTG TGG CAC 3' PCR amplification was performed using high fidelity

conditions (Eckert, K.A. and Kunkel, T.A. 1990 Nucl Acids 30 Research 18 3739-3744). The PCR mixture contained: 20mM TrisHCl (pH7.3 at 70°C, 50mM KCl, 4mM magnesium chloride, 0.01% gelatin, lmM each of dATP, dCTP, dGTP and dTTP, 500ng/ml RP41 DNA, luM each primer and 50 units/ml Tag polymerase (Cetus/Perkin Elmer, Beaconsfield, Bucks, 35 U.K.). Thirty cycles of PCR were performed with denaturation at 92°C for 1 min, annealing at 60°C for

lmin and extension at 72°C for 1.5 min. This reaction resulted in amplification of a fragment of ca. 1500bp as

expected. 40

fdCAT2 vector DNA (see example 5) was digested with ApaLl and Xhol (New England Biolabs) according to manufacturers recommendations, extracted with phenol/chloroform and ethanol precipitated (Sambrook et al. supra). Cloning of amplified RP41 DNA into this vector and identification of the desired clones was performed essentially as in example 7 except that digestion of the PCR product was with ApaLl and Xho 1. Colonies containing h-PDGFB-R DNA were identified by probing with 32p labelled RPDGF2 and the presence of an insert in hybridising colonies was confirmed by analytical PCR using RPDGF3 and RPDGF2 using the

PCT/GB91/01134

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conditions described in example 7.

Example 16

Binding of 1251-PDGF-BB to the Extracellular Domain of the Human Receptor for Platelet Derived Growth Factor Isoform BB Displayed on the Surface of fd Phage. Measured using an Immunoprecipitation Assay.

Phage particles, expressing the extracellular domain of the human platelet derived growth factor isoform BB receptor (fd h-PDGFB-R), were prepared by growing E.coli MCIO61 cells transformed with fd h-PDGFB-R in 50ml of 2xTY medium with 15ug/ml tetracycline for 16 to 20 hours. Phage particles were concentrated using polyethylene glycol as described in example 6 and resuspended in PDGF

glycol as described in example 6 and resuspended in PDGF binding buffer (25mM HEPES, pH7.4, o.15mM NaCl, lmM magnesium chloride, 0.25% BSA) to 1/337d of the original volume. Residuel bacteria and undissolved material were removed by spinning for 2 min in a mocrocentrifuge. Immunoblots using an antiserum raised against gene IIII

20 protein (Prof. I. Rashed, Konstanz, Germany) show the presence in such phage preparations of a geneIII-h-PDGFE-R protein of molecular mass 125000 corresponding to a fusion between h-PDGFB-R external domain (55000 daltons) and geneIII (apparent molecular mass 70000 on SDS-Dolvarylamide cel)

5 polyacrylamide gel). Duplicate samples of 35ul concentrated phage were incubated with 1251-PDGF-BB (78.7fmol, 70nCi, 882Ci/mmol; Amersham International plc, Amersham, Bucks) for 1 hour

at 37°C. Controls were included in which fdTPs/Hs vector phage (figure 4) or no phage replaced fd h-BDGFB-R phage. After this incubation, loul of sheep anti-M13 polyedonal antiserum (a gift from M. Hobart) was added and incubation continued for 30 min at 20°C. To each sample, 40ul (20ul packed volume) of protein G Sepharose Fast

35 Flow (Pharmacia, Milton Keynes) equilibrated in PDGF binding buffer was added. Incubation was continued for 30 min at 20°C with mixing by end over end inversion on a rotating mixer. The affinity matrix was spun down in a microcentrifuge for 2 min and the supernatant removed by

40 aspiration. Non-specifically bound ¹²⁵I-PDGF-BB was removed by resuspension of the pellet in 0.5ml PDGF binding buffer, mixing by rotation for 5 min. centrifugation and aspiration of the supernatant, followed by two further washes with 0.5ml 0.1% 85A, 0.2%

45 Triton-X-100. The pellet finally obtained was resuspended in 100ul PDGF binding buffer and counted in a Packard gamma counter. For displacement studies, unlabelled PDGF-BB (Amersham International) was added to the stated concentration for the incubation of 125I-PDGF-BB with phage.

 $1_{25}1\mbox{-PDGF-BB}$ bound to the fd h-PDGFB-R phage and was immunoprecipitated in this assay. Specific binding to

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receptor phage was 3.5 to 4 times higher than the nonspecific binding with vector phage fdTPs/Bs or no phage (fig. 19). This binding of $^{125}\text{I-PDGF-BB}$ could be displaced by the inclusion of unlabelled PDGF-BB in the incubation with phage at 37°C (fig. 20). At 50nM, unlabelled PDGF-BB the binding of $^{125}\text{I-PDGF-BB}$ was reduced to the same level as the fdTPs/Bs and no phage control. Figure 21 shows the same data, but with the non-specific binding to vector deducted.

These results indicate that a specific saturable site for $^{125}\text{I-PDGF-BB}$ is expressed on fd phage containing cloned h-PDGFB-R DNA. Thus, the phage can display the functional extracellular domain of a cell surface

receptor.

Example 17, Construction of Phagemid Containing GeneIII 15 fused with the Coding Sequence for a Binding Molecule It would be useful to improve the transfection

efficiency of the phage-binding molecule system and also to have the possibility of displaying different numbers and specificities of binding molecules on the surface of the same bacteriophage. The applicants have devised a

method that achieves both aims. The approach is derived from the phagemid system based on pUC119 [Vieira, J and Messing, J. (1987) Methods Enzymol. 153:3]. In brief, gene III from fd-CAT2 (example 5) and gene III scFv fusion from fd-CAT2 scFv 25 D1.3 (example 2) were cloned downstream of the lac promoter in separate samples of pUC119, in order that the inserted gene III and gene III fusion could be 'rescued' by M13M07 helper phage [Vieira, J and Messing, J. et 30 supra.) prepared according to Sambroots et al. 1989 supra. The majority of rescued phage would be expected to contain a genome derived from the pUC119 plasmid that contains the binding molecule-gene III fusion and should express varying numbers of the binding molecule on the 35 surface up to the normal maximum of 3-5 molecules of gene

III of the surface of wild type phage. The system has been exemplified below using an antibody as the binding molecule. An fdCAT2 containing the single chain Fv form of the 40 D1.3 antilysozyme antibody was formed by digesting

fdTscFvD1.3 (example 2) with Pstl and Xhol, purifying the fragment containing the scFv fragment and ligating this into Pstl and Xhol digested fdCAT2. The appropriate clone, called fdCAT2 scFvD1.3 was selected after plating 45 onto 2xTY tetracycline (15µg/ml) and confirmed by

restriction enzyme and sequence analysis.

Gene III from fd-CAT2 (example 5) and the gene III scPv fusion from fd-CAT2 scFvDl.3 was PCR-amplified using the primers A and B shown below:

Primer A: TGC GAA GCT TTG GAG CCT TTT TTT TTG GAG ATT TTC AAC G

Primer E: CAG TGA ATT CCT ATT AAG ACT CCT TAT TAC GCA GTA TGT TAG C

Primer A anneals to the 5' end of gene III including the ribosome binding site is located and incorporates a Hind III site. Primer B anneals to the 3' end of gene III at the C-terminus and incorporates two UAA stop codons and an EcoRI site. 100 ng of fd-CAT2 and fd-CAT2 scFv Dl.3 DNA was used as templates for PCR-amplification in a total reaction volume of 50µl as described in example 7, except that 20 cycles of amplification were performed: 94°C 1 minute, 50°C 1 minute, 72°C 3 minutes. This resulted in amplification of the expected 1.2Kb fragment from fd-CAT2 and a 1.8Kb fragment from fd-CAT2

scFv Dl.3.

The PCR fragments were digested with EcoR1 and Hind III, gel-purified and ligated into Eco-R1- and Hind III-cut and dephosphorylated pUCl19 DNA and transformed into E.coli TG1 using standard techniques (Sambrook et al., et supra). Transformed cells were plated on SOB agar (Sambrook et al. 1989 supra) containing 100µg/ml ampicillm and 2% glucose. The resulting clones were called pCAT-3 (derived from fd-CAT2) and pCAT-3 scFv Dl.3 (derived from fd-CAT2 scFv Dl.3).

Example 18, Rescue of Anti-Lysozyme Antibody Specificity
75 from pCAT-3 scry D1 3 by M13807

from pCAT-3 scFv Dl.3 by Ml3KO7 Single pCAT-3 and pCAT-3 scFv Dl.3 colonies were picked into 1.5ml 2TY containing 100µg/ml ampicillin and 2% glucose, and grown 6 hrs at 30°C. 30µl of these stationary cells were added to 6mls 2YT containing 100ug/ml ampicillin and 2% glucose in 50ml polypropylene 30 tubes (Falcon, Becton Dickinson Labware, 1950 Williams Drive, Oxnard, CA. USA) and grown for 1.5 hrs at 30°C at 380rpm in a New Brunswick Orbital Shaker (New Brunswick Scientific Ltd., Edison House 163 Dixons Hill road, North 35 Mimms, Hatfield, UK). Cells were pelleted by centrifugation at 5,000g for 25 minutes and the tubes drained on tissue paper. The cell pellets were then suspended in 6mls 2TY containing 1.25x109 p.f.u. ml-1 ML3KO7 bacteriophage added. The mixture was left on ice 40 for 5 minutes followed by growth at 35°C for 45 minutes at 450rpm. A cocktail was then added containing 4ul 100µg/ml ampicillin, 0.5µl 0.1M TPTG and 50µl 10mg/ml kanamycin, and the cultures grown overnight at 35°C. 450rpm.

The following day the cultures were centrifuged and phage particles PEG precipitated as described in example 6. Phage pellets were resuspended in 100µl TE (tris-EDTA see example 6) and phage titred on E.coli TGL. Aliquots of infected cells were plated on 2TV containing either 100µg/ml ampicillin to select for pUC119 phage particles, or 50µg/ml kanamycin to select for the M13 KO7 heiper phage. Plates were incubated overnight at 37°C and

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antibiotic-resistant colonies counted: ampR

kanR DNA 1.8x10¹¹ colonies 1.2x10⁹ colonies pCAT-3

2.4x10¹¹ colonies 2.0x10⁹ colonies DCAT-3scFv D1.3 This shows that the ampR phagemid particles are infective and present in the rescued phage population at

a 100-fold excess over kanR M13K07 helper phage. Phage were assayed for anti-lysozyme activity by

ELISA as described in example 6, with the following modifications: 10

ELISA plates were blocked for 3 hrs with 2% Marvel/PBS.

50µl phage, 400µl 1xPBS and 50µl 20% Marvel were mixed end over end for 20 minutes at room temperature before adding 150µl per well.

Phage were left to bind for 2 hours at room

temperature.

All washes post phage binding were: 2 quick rinses PBS/0.5% Tween 20

3x2 minute washes PBS/0.5% Tween 20 2 quick rinses PBS no detergent 3x2 minute washes PBS no detergent

The result of this ELISA is shown in figure 22, which shows that the antibody specificity can indeed be rescued efficiently.

It is considered a truism of bacterial genetics that when mutant and wild-type proteins are co-expressed in the same cell, the wild-type proteins are co-expressed in same cell, the wild-type protein is used preferentially. This is analogous to the above situation wherein mutant (i.e. antibody fusion) and wild-type gene III proteins (from M13K07) are competing for assembly as part of the pUC119 phagemid particle. It is therefore envisaged that the majority of the resulting pUC 119 phage particles will have fewer gene III-antibody fusion molecules on their surface than is the case for purely phage system described for instance in example 2. Such phagemid antibodies are therefore likely to bind antigen with a

lower avidity than fd phage antibodies with three or more copies of the antibody fusion on their surfaces (there is 40 no wild-type gene III, in the system described, for instance, in example 2), and provide a route to production of phage particles with different numbers of the same binding molecule (and hence different acidities for the ligand/antigen) or multiple different binding

specificities on their surface, by using helper phage such as M13K07 to rescue cells expressing two or more gene III-antibody fusions.

It is also possible to derive helper phage that do not encode a functional gene III in their genomes (by for 50 example deleting the gene III sequence or a portion of it or by incorporating an amber mutation within the gene).

These defective phages will only grow on appropriate cells (for example that provide functional gene III in trans, or contain an amber supressor gene), but when used to rescue phage antibodies, will only incorporate the gene III antibody fusion encoded by the phagemid into the released phage particle.

Example 19. Transformation Efficiency of pCAT-3 and

pCAT-3 scFv Dl.3 phagemids

pUC 19, pCAT-3 and pCAT-3 scFv D1.3 plasmid DNAs. 10 and fdCAT-2 phage DNA was prepared, and used to transform E.coli TG1, pCAT-3 and pCAT-3 scFv D1.3 transformations were plated on SOB agar containing 100µg/ml ampicillin and 2% glucose, and incubated overnight at 30°C. fdCAT-2 transformations were plated on TY agar containing 15µg/ml 15 tetracycline and incubated overnight at 37°C.

Transformation efficiencies are expressed as colonies per

ug of input DNA.

	DNA	Transformation	efficiency
20	pUC 19	1.109	
	pCAT-3	1.108	
	pCAT-3scFv Dl.3	1.108	
	fd CAT-2	8.105	
	X n numman A	the second common de de common	

As expected, transformation of the phagemid vector 25 is approximately 100-fold more efficient that the parental fdCAT-2 vector. Furthermore, the presence of a SCFV antibody fragment does not compromise efficiency. This improvement in transformation efficiency is practically useful in the generation of phage antibodies 30

libraries that have large repertoires of different binding specificities.

Example 20

PCR Assembly of a Single Chain Fv Library from an

Immunised Mouse

To demonstrate the utility of phage for the selection of 35 antibodies from repertoires, the first requirement is to be able to prepare a diverse, representative library of the antibody repertoire of an animal and display this repertoire on the surface of bacteriophage fd.

40 Cytoplasmic RNA was isolated according to example 14

from the pooled spleens of five male Balb/c mice boosted 8 weeks after primary immunisation with 2-phenyl-5oxazolone (ph OX) coupled to chicken serum albumin. CDNA preparation and PCR assembly of the mouse VH and VL kappa 45 repertoires for phage display was as described in example The molecules thus obtained were ligated into

fdCAT2.

Vector fdCAT2 was extensively digested with Notl and ApaLl., purified by electroelution (Sambrook et al.a989 50 supra) and 1 µg ligated to 0.5 µg (5 µg for the hierarchial libraries: see example 22) of the assembled scFv genes in 1 ml with 8000 units T4 DNA ligase (New

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England Biolabs). The ligation was carried out overnight at 16°C. Purified ligation mix was electroporated in six aliquots into MC1061 cells (W. J. Dower, J. F. Miller & C. W. Ragsdale Nucleic Acids Res. 16 6127-6145 1988) and plated on NZY medium (Sambrook et al. 1989 supra) with 15pg/ml tetracycline, in 243x243 mm dishes (Nunc): 90-95%

of clones contained scrv genes by PCR screening.

Recombinant colonies were screened by PCR (conditions as

in example 7 using primers VHIBACK and MJKIFONX, MJK2FONX, MJK4FONX and MJK5FONX (see example 14) followed 10 by digestion with the frequent cutting enzyme BstN1 (New England Biolabs, used according to the manufacturers instructions). The library of 2x105 clones appeared diverse as judged by the variety of digestion patterns seen in Figure 23, and sequencing revealed the presence 15

of most VH groups (R. Dildrop, Immunol. Today 5 85-86. 1984) and VK subgroups (Kabat. E.A. et al. 1987 supra) (data not shown). None of the 568 clones tested bound to

phOx as detected by ELISA as in example 9.

Thus the ability to select antibody provided by the 20 use of phage antibodies (as in example 21) is essential to readily isolate antibodies with antigen binding activity from randomly combined VH and VL domains. Very extensive screening would be required to isolate antigenbinding fragments if the random combinatorial approach of 25

Huse et al. 1989 (supra) were used.

Example 21

Selection of Antibodies Specific for 2-phenyl-5-oxazolone from a Repertoire Derived from an Immunised Mouse
The library prepared in example 20 was used to

demonstrate that ability of the phage system to select antibodies on the basis of their antibody specificity.

None of the 568 clones tested from the unselected

library bound to phOx as detected by ELISA.

Screening for binding of the phage to hapten was carried out by ELISA: 96-well plates were coated with 10 µg/ml phOx-BSA or 10 µg/ml BSA in phosphate-buffered saline (PBS) overnight at room temperature. Colonies of phage-transduced bacteria were inoculated into 200 µl 2 x

TY with 12.5 µg/ml tetracycline in 95-well plates ('cell wells', Nuclon) and grown with shaking (300 rpm) for 24 hours at 37°C. At this stage cultures were saturated and phage titres were reproducible (10¹⁰ TU/m1). 50 µl phage supernatant, mixed with 50 µl PBS containing 4% skimmed milk powder, was then added to the coated plates.

45 Further details as in example 9.

The library of phages was passed down a phOx affinity column (Table 4A), and eluted with hapten. Colonies from the library prepared in example 22 were scraped into 50ml 2 x TY medium37 and shaken at 37°C for 50 Liberated phage were precipitated twice with polyethylene glycol and resuspended to 1012 TU

(transducing units)/ml in water (titred as in example 8). For affinity selection, a 1 ml column of phOx-BSA-Sepharose (O. Makela, M. Kaartinen, J.L.T. Pelonen and K. Karjalainen J. Exp. Med. 148 1644-1660, 1978) was washed with 300 ml phosphate-buffered saline (PBS), and 20 ml PBS containing 2% skimmed milk powder (MPBS). 1012 TU phage were loaded in 10 ml MPBS, washed with 10 ml MPBS and finally 200 ml PBS. The bound phage were eluted with 5 ml 1 mM 4-6-amino-caproic acid methylene 2-phenyl-Oxazol-5-one phox-CAP; O. Makela et al. 1978, supra). About 10° TU eluted phage were amplified by infecting 1 ml log phase E.coli TGl and plating as above. For a 10 further round of selection, colonies were scraped into 10 ml 2 x TY medium and then processed as above. Of the eluted clones, 13% were found to bind to phOx after the

15 first round selection, and ranged from poor to strong binding in ELISA.

To sequence clones, template DNA was prepared from the supernatants of 10 ml cultures grown for 24 hours. 20 and sequenced using the dideoxy method and a Sequenase kit (USB), with primer LINKFOR (see example 14) for the VH genes and primer fdSEQ1 (5'-GAA TTT TCT GTA TGA GG) for the Vk genes. Twenty-three of these hapten-binding clones were sequenced and eight different VH genes (A to 25 H) were found in a variety of pairings with seven different Vk genes (a to g) (Fig. 24). Most of the domains, such as VH-B and Vk-d were 'promiscuous', able to bind hapten with any of several partners.

The sequences of the V-genes were related to those 30 seen in the secondary response to phOx, but with differences (Fig. 24). Thus phOx hybridomas from the secondary response employ sometically mutated derivatives

of three types of Vk genes - Vkoxl. t'Vkox-like' and Vk45.1 genes (C. Berek, G. M. Griffiths & C. Milstein Nature 316 412-418 (1985). These can pair with VH genes 35 from several groups, from Vkoxl more commonly pairs with the VHoxl gene (VH group 2. R.Dildrop uupra). genes are always, and Vkox-like genes often, found in association with heavy chains (including VHox1) and

contain a short five residue CDR3, with the sequence motif Asp-X-Gly-X-X in which the central glycine is 40 needed to create a cavity for phOx. In the random combinatorial library however, nearly all of the VH genes belonged to group 1, and most of the Vk genes were ox-

45 like and associated with VH domains with a five residue CDR3, motif Asp/Asn-X-Gly-X-X (Fig. 24). Vkoxl and VHoxl were found only once (Vk-f and VH-E), and not in combination with each other. Indeed Vk-f lacks the Tro91 involved in phOx binding and was paired with a VH (VH-C) 50

with a six residue CDR3.

A matrix combination of VH and VK genes was identified in phOx-binding clones selected from this

random combinational library. The number of clones found with each combination are shown in Fig. 25. The binding to phOx-BSA, as judged by the ELISA signal, appeared to vary (marked by shading in Fig. 25). No binding was seen to BSA alone.

A second round of selection of the original, random combinational library from immune mice resulted in 93% of eluted clones binding phOx (Table 4). Most of these clones were Vk-d combinations, and bound strongly to phox in ELISA (data not shown). Few weak binders were seen. This suggested that affinity chromatography had not only

enriched for binders, but also for the best.

Plorescence quench titrations determined the Kd of VH-B/Vk-d for phOx-GABA as 10-8 M (example 23). indicating that antibodies with affinities representative 15 of the secondary response can be selected from secondary response, only two (out of eleven characterised) secrete antibodies of a higher affinity than VH-B/Vk-d (C. Berek et al. 1985 supra). The Kd of VH-B/Vk-b for phOx-GABA was determined as 10-5 M (example 23). Thus phage 20 bearing scFv fragments with weak affinities can be

selected with antigen, probably due to the avidity of the

nultiple antibody heads on the phage.

This example shows that antigen specificities can be isolated from libraries derived from immunised mice. It 25 will often be desired to express these antibodies in a soluble form for further study and for use in therapeutic Example 23 demonstrates and diagnostic applications. determination of the affinity of soluble scFv fragments selected using phage antibodies. Example 27 demonstrates 30 that soluble fragments have similar properties to those displayed on phage. For many purposes it will be desired to construct and express an antibody molecule which contains the Fc portions of the heavy chain, and perhaps vary the immunoglobulin isotype. To accomplish this, it 35

is necessary to subclone the antigen binding sites identified using the phage selection system into a vector for expression in mammalian cells, using methodology similar to that described by Orlandi, R. et al. (1989, supra). For instance, the VH and VL genes could be 40

amplified separately by PCR with primers containing appropriate restriction sites and inserted into vectors such as pSV-gpt HulgGl (L. Riechmann et al Nature 332 323-327), 1988) which allows expression of the VH domain as part of a heavy chain IgGl isotype and pSV-hyg HuCK

45 which allows expression of the VL domain attached to the K light chain constant region. Furthermore, fusions of VH and VL domains can be made with genes encoding nonimmunoglobulin proteins, for example, enzymes.

50 Example 22

Generation of Further Antibody Specificities by the Assembly of Hierarchical Libraries

Further antibody specificities were derived from the library prepared and screened in examples 20 and 21 using a hierarchical approach.

The promiscuity of the VH-B and Vk-d domains prompted the applicants to force further pairings, by assembling these genes with the entire repertoires if either Vk or VH genes from the same immunised mice. The resulting 'hierarchical' libraries, (VH-B x Vk-rep and VH-rep x Vk-d), each with 4×10^7 members, were subjected to a round of selection and hapten-binding clones isolated (Table 4). As shown by ELISA, most were strong binders. By sequencing twenty-four clones from each library, the applicants identified fourteen new partners for VH-B and thirteen for Vk-d (Fig. 24). Apart from VH-

B and Vk-c, none of the previous partners (or indeed other clones) from the random combinatorial library was 15 isolated again. Again the Vk genes were mainly ox-like and the VH genes mainly group I (as defined in Dildrop, R. 1984 supra), but the only examples of Vkoxl (Vk-h. -D. 20 -q and -r) have Trp91, and the VH-CDR3 motif Asp-X-Gly-X-

X now predominates. Thus some features of the phOx hybridomas seemed to emerge more strongly in the hierarchial library. The new partners differed from each other mainly by small alterations in the CDRs, indicating 25 that much of the subtle diversity had remained untapped by the random combinatorial approach. More generally it has been shown that a spectrum of related antibodies can be made by keeping one of the partners fixed and varying the other, and this could prove invaluable for fine

30 tuning of antibody affinity and specificity. Therefore, again, phage antibodies allow a greater range of antibody molecules to be analysed for desired

properties.

This example, and example 21, demonstrate the 35 isolation of individual antibody specificities through display on the surface of phage. However, for some purposes it may be more desirable to have a mixture of antibodies, equivalent to a polyclonal antiserum (for instance, for immunoprecipitation). To prepare a mixture 40 of antibodies, one could mix clones and express soluble antibodies or antibody fragments or alternatively select clones from a library to give a highly enriched pool of genes encoding antibodies or antibody fragments directed against a ligand of interest and express antibodies from

45 these clones. Example 23

Selection of Antibodies Displayed on Bacteriophage with Different Affinities for 2-phenyl-5-oxezolone using Affinity Chromatography

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The ELISA data shown in example 21 suggested that affinity chromatography had not only enriched for binders, but also for the best. To confirm this, the

binding affinities of a strong binding and a weak binding phage were determined and then demonstrated that they could be separated from each other using affinity chromatography.

chromatography.

Clones VH-B/VK-b and VH-B/VK-d were reamplified with MJK1FONX, MJK2FONX, MJK4FONX and MJK5FONX (see example 14) and VHLBACK-Sfil (5'-TCG CGG CCC AGC CGG CCA TGG CC(G/C) AGG T(C/G)(A/C) A(A/G)C TGC AG(C/G) AGT C(A/T)G G), a primer that introduces an Sfil site (underlined) at the 5' end of the VH gene. VH-B/VK-d was cloned into a phagemid e.g. pJM1 (a gift from A. Griffiths and J. Marks) as an Sfil-Notl cassette, downstream of the pelB leader for periplasmic secretion (M. Better at al. supra), with a C-terminal peptide tag for detection (see example 24 and figure), and under th control of a Fg.

example 24 and figure), and under th control of a Promoter (H. Shimatake & M. Rosenberg Nature 292 128-132 1981). The phagemid should have the following features:

a) unique Sfil and Notl restriction sites downstream of a pelB leader; b) a sequence encoding a C-terminal peptide

20 tag for detection; and c) a / P, promoter controlling expression. 10 litre cultures of E.coli N4830-1 (M. E. Gottesman, S. Adhya & A. Das J.Mol.Biol 140 57-75 1980) harbouring each phagemid were induced as in K. Nagai & H. C. Thogerson (Methods Enzymol 153 461-481 1987) and supernatants precipitated with 50% ammonium sulphate. The resuspended precipitate was dialysed into PBS + 0.2

mM EDTA (PBSE), loaded onto a 1.5ml column of phOx:Sepharose and the column washed sequentially with 100 ml PBS: 100 ml 0.1 M Tris-HCl, 0.5 M NaCl, pH 8.0: 30 10ml 50 mM citrate, pH 5.0: 10 ml 50 mM citrate, pH4.0, and 20 ml 50 mM glycine, pH 3.0. scFv fragments were

eluted with 50 mM glycine, pH 2.0, neutralised with Tris base and dialysed against PBSE. VH-B/Vk-b was cloned into a phagemid vector based on pUCl19 encoding identical signal and tag sequences to pJM1, and expression induced at 30°C in a 10 litre culture of E.coli TG1 harbouring

the phagemid as in D. de Bellis & I. Schwartz (1980 Nucleic Acids Res 18 131). The low affinity of clone VH-B/VK-b made its purification on phOx-Sepharose do impossible. Therefore after concentration by ultrafiltration (Filtron, Flowgen), the supernatant (100

ml of 600 ml) was loaded onto a 1 ml column of protein A-Sepharose cpoupled (E. Harlow & D. Lane 1988 supra) to the monoclonal antibody 9E10 (Evan, G. I. et al. Mol.Cell Biol.5 3610-3616 1985) that recognises the peptide tag. The column was washed with 200 ml PBS and 50 ml PBS made

The column was washed with 200 ml PBS and 50 ml PBS made 0.5 M in NaCl. scPv fragments were eluted with 100 ml 0.2M glycine, pH 3.0, with neutralisation and dialysis as before.

The Kd (1.0 + 0.2 x 10⁻⁸ M) for clone VH-B/Vk-d was

The Kd (1.0 + 0.2 x 10⁻⁸ M) for clone VH-B/Vk-d was determined by fluorescence quench titration with 4-E-amino-butyric acid methylene 2-phenyl-oxazol-5-one (phox-

GABA Co. Makela et al. 1978 supra). Excitation was at 280 nm, emission was monitored at 340 nm and the $\rm K_{\rm d}$ calculated. The $\rm K_{\rm d}$ of the low affinity clone VH-B/VK-b was determined as $1.8\pm~0.3\times10^{-5}$ M (not shown). To minimise light adsorption by the higher concentrations of phOx-GABA required, excitation was at 260 nm and emission was monitored at 304 nm. In addition the fluorescence values were divided by those from a parallel titration of the lysozyme binding Fv fragment D1.3. The value was calculated as in H. N. Eisen Meth.Med.Res. 10 115-121 1964. A mixture of clones VH-B/Vk-b and VH-B/Vk-d, 7x10 10 TU phage in the ratio 20 VH-B/Vk-b: 1 VH-B/VK-d were loaded onto a phOx-BSA-Sepharose column in 10 ml MPBS and eluted as above. Eluced phage were used to

15 reinfect E.coli TGI, and phage produced and harvested as before. Approximately 10¹¹ TU phage were loaded onto a second affinity column and the process repeated to give a total of three column passes. Dilutions of eluted phage at each stage were plated in duplicate and probed separately with cligonuclectides specific for Vk-b (5'GAC CGG GTA ACC ACT GTA CT) or Vk-d (5'-GAA TGG TAT AGT ACT

ACC CT). After these two rounds, essentially all the eluted phage were VH-B/Vk-d (table 4). Therefore phage antibodies can be selected on the basis of the antigen affinity of the antibody displayed.

Example 24

Construction of Phagemid pHEN1 for the Expression of Antibody Fragments Expressed on the Surface of Bacteriophage following Superinfection

30 The phagemid pHEN1 (figure 26) is a derivative of

pUCL19 (Vieire, J. & Messing, J. Methods Enzymol 153 pp 3-11, 1987). The coding region of g3p from fdCAT2, including signal peptide and cloning sites, was amplified by FCR, using primers G3FUFO and G3FUBA (given below) (which contain EcoRI and HindIII sites respectively), and cloned as a HindIII-EcoRI fragment into pUCl19. The HindIII-NotI fragment encoding the g3p signal sequence was the replaced by a pelB signal peptide (Better, M. et al. Science 240 1041-1043, 1988) with an internal SfiI site, allowing antibody genes to be cloned as fiI-NotI fragments. A peptide tag, c-myc, (Munro, S. & Pelham, H. Call 46 291-300, 1986) was introduced directly after the

NotI site by cloning an oligonucleotide cassette, and followed by an amber codon introduced by site-directed mutagenesis using an in vitro mutagenesis kit (Amersham International) (figure 26b).

GSFUFO, 5'-CAG TGA ATT CTT ATT AAG ACT CCT TAT TAC GCA GTA TGT TAG C; GSFUFA, 5'-TGC GAA GCT TTG GAG CCT TTT TTT TTG GAG ATT TTC

AAC G:

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Example 25 Display of Single Chain Fy and Fab Fragments Derived from the Anti-Oxazolone Antibody NQ10.12.5 on Bacteriophage fd

using pHEN1 and fdCAT2

A range of constructs (see figure 27) were made from a clone (essentially construct II in pUC19) designed for expression in bacteria of a soluble Fab fragment (Better et al. 1988 see above) from the mouse anti-phOx (2phenyl-5-oxazolone) antibody NQ10.12.5 (Griffiths, G. M. et al. Nature 312, 271-275, 1984). In construct II, the V-regions are derived from NQ10.12.5 and attached to human Ck and CH1 (%1 isotype) constant domains. terminal cysteine residues, which normally form a

covalent link between light and heavy antibody chains, 15 have been deleted from both the constant domains. clone heavy and light chain genes together as Fab fragments (construct II) or as separate chains (constructs III and IV) for phage display, DNA was

amplified from construct II by PCR to introduce a NotI 20 restriction site at the 3' end, and at the 5' end either an ApaLIssite (for cloning into fd-CAT2) or Sfil sie (for into pHEN1). The primers FABNOTFOK with cloning VHIBACKAPA (or VHIBACKSFI15) were used for PCR

amplification of genes encoding Fab fragments (construct 25 II), the primers FABNOTFOH with VHIBACKAPA (or VHIBACKSFILS) for heavy chains (construct III), and the primers FABNOTFOK and MVKBAAPA (or MVKBASFI) for light chains (construct IV).

The single-chain Fv version of NQ10.12.5 (construct 30 I) has the heavy (VH) and light chain (Vk) variable domains joined by a flexible linker (Gly4Ser) (Huston, J. S. et al. Proc. Natl. Acad. Sci. USA 85 5879-5883, 1988) and was constructed from construct II by 'splicing

by overlap extension' as in example 14. The assembled 35 genes were reamplified with primers VK3F2NOT and VHIBACKAPA (or VHIBACKSFI15) to append restriction sites for cloning into fd-CAT2 (ApaLI-NotI) or pHEN1 (Sfil-

NotI). 40

VHIBACKAPA, 5'-CAT GAC CAC AGT GCA CAG GT(C/G) (A/C)A(A/G) CTG CAG (C/G)AG TC(A/T) GG; VHIBACKSFI15,5'-CAT GCC ATG ACT CGC GGC CCA GCC GGC CAT GGC C(C/G)A GGT (C/G)(A/C)A (A/G)CT GCA G(C/G)A GTC

45 (A/T)GG; FABNOTFOH, 5'-CCA CGA TTC TGC GGC CGC TGA AGA TTT GGG CTC AAC TTT CTT GTC GAC; FABNOTFOK, 5'-CCA CGA TTC TGC GGC CGC TGA CTC TCC GCG GTT GAA GCT CTT TGT GAC;

MYKBAAPA, 5'-CAC AGT GCA CTC GAC ATT GAG CTC ACC CAG TCT 50 MVKBASFI,5'-CAT GAC CAC GCG GCC CAG CCG GCC ATG GCC GAC WO 92/01047 PCT/GB91/01134

81

ATT GAG CTC ACC CAG TCT CCA; VK3F2NOT,5'-TTC TGC GGC CGC CCG TTT CAG CTC GAG CTT GGT CCC.

Restriction sites are underlined.

Rescue of Phage and Phagemid particles
Constructs I-IV (figure 27) were introduced into both fd-

CAT2 and pHEN1. Phage 6d-CAT2 (and fd-CAT2-I,II,III or IV) was taken from the supernatant of infected E.coli TG1 after shaking at 37°C overnight in 2xTY medium with 10 12.5µg/ml tetracycline, and used directly in ELISA. Phagemid pHEN1 (and pHEN1-I and II) in E.coli TG1 (supE) were grown overnight in 2 ml 2xTY medium, 100 µg/ml ampicillin, and 1% glucose (without glucose, expression of g3p prevents later superinfection by helper phage).

15 10µl of the overnight culture was used to innoculate 2 ml of 2xTY medium, 100µg/ml ampicillin, 1% glucose, and shaken at 37°C for 1 hour. The cells were washed and resuspended in 2xTY, 100 µg/ml ampicillin, and aphagemid particles rescued by adding 2 µl (108pfu) VCSM13 helper phage (Stratagene). After growth for one hour, 4µl kanamycin (25 mg/ml) was added, and the culture grown overnight. The phagemid particles were concentrated 10-

kanamycin (25 mg/ml) was added, and the culture grown overnight. The phagemid particles were concentrated 10fold for ELISA by precipitation with polyethylene glycol. ELISA Detection of phage binding to 2-phenyl-5-oxazolone (phOx)

25 Detection of phage binding to 2-phenyl-5-oxazolone (phOx) was performed as in example 9. 96-well plates were coated with 10 µg/ml phOx-BA or 10 µg/ml BSA in PBS overnight at room temperature, and blocked with PBSS containing 2% skimmed milk powder. Phage (mid) 30 supernatant (50 µl) mixed with 50 µl PBS containing 4% skimmed milk powder was added to the wells and assayed. To detect binding of soluble scFv or Fab fragments secreted from plEBNl, the c-myc peptide tag described by Munro and Pelham 1986 supra, was detected using the anti-

35 myc monoclonal 9E10 (Evan, G. I. et al. Mol Cell Biol 5 3610-3616, 1985) followed by detection with peroxidaseconjugated goat anti-mouse immonoglobulin. Other details are as in example 9.

The constructs in fdCATZ and pHEN1 display antibody
fragments of the surface of filamentous phage. The phage
vector, fd-CATZ (figure 8) is based on the vector fd-tet
(Zacher, A. N. et al. Gene 9 127-140, 1980) and has
restriction sites (ApaLI and Not1) for cloning antibody
genes (or other protein) genes for expression as fusions
to the N-terminus of the phage coat protein gap.
Transcription of the antibody-gap fusions in fd-CATZ is
driven from the gene III promoter and the fusion protein
targetted to the periplasm by means of the g3p leader.
Fab abd scFv fragments of NQ10.12.5 cloned into fd-CATZ
for display were shown to bind to phOx-BSA (but not BSA)
by ELISA (table 5). Phage were considered to be binding

if Agos of the sample was at least 10-fold greater that

the background in ELISA.

The phagemid vector, pHENI (fig. 26), is based upon pUCl19 and contains restriction sites (SfiI and NotI) for cloning the fusion proteins. Here the transcription of antibody-g3p fusions is driven from the inducible lact promoter and the fusion protein targetted to the periplasm by means of the pelB leader. Phagemid was rescued with VCSMI3 helper phage in ZxTY medium containing no glucose or IFTG: under these conditions there is sufficient expression of antibody-g3p. Fab and scFy fragments of NOIO.12.5 cloned into pHENI for display

there is sufficient expression of antibody-gap. Fab and scPv fragments of NOIO.12.5 cloned into pHENI for display were shown to bind to phOx-BSA (but not BSA) by ELISA (Table 5) using the same criterion as above. An alternative methodology for preparing libraries

15 of Fab fragments expressed on the surface of phage would be to:

1. Prepare a library of phage expressing heavy chain (VHCH) genes from inserts in the phage genome.

 Prepare a library of light chain genes in a plamid expression vector in E.coli, preferably a phagemid, and isolate the soluble protein light chins expresed from this library.

 Bind the soluble protein light chains fromt he library to the heavy chain library displayed on phage.

25 4. Select phage with the desired properties of affinity and specificity.

These will encode the heavy chain (VHCH) genes.

5. Isolate the light chain genes encoding ight chains which form suitable antigen binding sites in combination with the selected heavy chains, preferably by using superinfectin of bacteria, containing phagemid expressing the light chain, with phage expressing the selected heavy chain (as described in example 20) and then assaying for antigen binding.

SEXEMPLE 26
RESCUE OF Phagemid Encoding a Gene III Protein Fusion
With Antibody Reavy or Light Chains by Phage Encoding the
Complementary Antibody Chain Displayed on Phage and the
Use of this Technique to Make Dual Combinatorial

40 Libraries
With random combinatorial libraries there is a
limitation on the potential diversity of displayed Fab
fragments due to the transformation efficiency of
bacterial cells. Described here is a strategy (dual

45 combinatorial libraries) to overcome this problem, potentially increasing the number of phage surveyed by a factor of 10'.

Por assembly of heavy and light chains expresses from different vectors, phagemid (pHEN1-III or IV) was grown in E.coli HB2151 (a non-supressor strain) to allow production of soluble chains, and rescued as above (example 27) except that helper phage were used

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expressing partner chains as fusions to g3p (10 9 TU fd-CAT2-IV or III respectively) and 2 μ l tetracycline (12.5 mg/ml) in place of kanamycin.

Separate Vectors to Encode Fab Heavy and Light Chains

The heavy and light chains of Fab fragments can be encoded together in the same vector (example 25) or in different vectors. To demonstrate this the heavy chain (construct III) was cloned into pHEN1 (to provide soluble fragments) and the light chain (construct IV) into fd-CAT2 (to make the fusion with g3p). The phagemid pHEN1-III, grown in E.ooli HB2151 (non-supressor) was rescued with fd-CAT2-IV phage, and phage(mid) shown to bind to phOx:BSA, but not to BSA (Table 5). This demonstrates that soluble light chain is correctly associating with the heavy chain anchored to the g3p, since neither heavy chain nor light chain lose bind antigen (Table 5).

Similar results were obtained in the reverse experiment (with phagemid pHEN-I-IV and fd-CAT2-III phage) in which the heavy chain was produced as a soluble molecule and the light chain anchored to g3p (Table 5). Hence a Fab fragment is assembled on the surface of phage by fusion of either heavy or light chain to g3p, provided the other chain is secreted using the same or another

25 vector (figure 28).

The resulting phage population is a mixture of phage abd rescued phagemid. The ratio of the two types of particle was assessed by infecting log phase E.coli TG1 and plating on TYE plates with either 15 µg/ml tetracycline (to select for fd-CAT2) or 100 µg/ml ampicillin (to select for pHEN1). The titre of fd-CAT2 phage was 5 x 101 TU/ml and the titre of pHEN1 2 x 1010 TU/ml, indicating a packaging ratio of 25 phage per phagemid.

35 Demonstrated here is an alternative strategy involving display of the heterodimeric antibody Fab fragments on the surface of phage. One of the chains is fused to g3p and the other is secreted in soluble form into the periplasmic space of the E.coli where it associates non-covalently with the g3p fusion, and binds specifically to antigen. Either the light or heavy chain can be fused to the g3p: they are displayed on the phage 40 as Fab fragments and bind antigen (Figure 28). Described are both phage and phagemid vectors for surface display. 45 Phagemids are probably superior to phage vectors for creation of large phage display libraries. Particularly in view of their higher transfection efficiencies (Two to three orders of magnitude higher), allowing larger libraries to be constructed. The phagemid vector, pHEN1 50 also allows the expression of soluble Fab fragments in non-suppressor E.coli.

Also demonstrated here is that heavy and light

chains encoded on the same vector (construct II), or on different vectors (constructs III and IV) can be This offers two distinct displayed as Fab fragments. ways of making random combinatorial libraries for Libraries of heavy and light chain genes, display. amplified by PCR, could be randomly linked by a 'PCR assembly' process (example 14) based on 'splicing by overlap extension', cloned into phage(mid) display vectors and expressed from the same promoter as part of the same transcript (construct II) as above, or indeed 10 from different promoters as separate transcripts. the phage(mid) vector encodes and_displays both chains, For a combinatorial library of 107 heavy chains and 107 light chains, the potential diversity of displayed Fab fragments (10^{14}) is limited by the transfection 15 efficiency of bacterial cells by the vector (about 109 clones per ug cut and ligated plasmid at best) (W.J. Dower et al Nucl. Acids. Res. 16 6127-6145, 1988). Libraries thus prepared are analogous to the random combinatorial library method described by Huse, W.D. et 20 al Science 246 1275-1281 (1989), but have the important additional feature that display on the surface of phage gives a powerful method of selecting antibody specificities from the large number of clones generated. Alternatively, libraries of heavy and light chains 25 could be cloned into different vectors for expression in the same cell, with a phage vector encoding the g3p fusion and a phagemid encoding the soluble chain. The phage acts as a helper, and the infected bacteria produced both packaged phage and phagemid. Each phage or 30 phagemid displays both chains but encodes only one chain and thus only the genetic information for half of the antigen-binding site. However, the genes for both antibody chains can be recovered separately by plating on the selective medium, suggesting a means by which 35 mutually complementary pairs of antigen binding heavy and light chain combinations could be selected from random For example, a light chain combinatorial libraries. repertoire on fd phage could be used to infect cells harbouring a library of soluble heavy chains on the 40 phagemid. The affinity purified phagemid library could then be used to infect E.coli, rescued with the affinity purified phage library, and the new combinatorial library subjected to a further round of selection. antibody heavy and light chain genes are reshuffled after 45 each round of purification. Finally, after several rounds, infected bacteria could be plated and screened individually for antigen-binding phage. Such 'dual' combinatorial libraries are potentially more diverse than those encoded on a single vector. By combining separate libraries of 10^7 light chain phage(mid)s, the diversity of displayed Fab fragments (potentially 10^{14}) is limited 50

only by the number of bacteria $(10^{12} \ \mathrm{per}\ \mathrm{litre})$. More simply, the use of two vectors should also facilitate the construction of 'hierarchical' libraries, in which a fixed heavy or light chain is paired with a library or partners (example 22), offering a means of 'fine-tuning' antibody affinity and specificity. Example 27

Induction of Soluble scFv and Fab Fragments using Phagemid DHENI

Further study of antibodies which have been expressed on the surface of phage would be greatly facilitated if it is simple to switch to expression in solution.

E.coli HE2151 was infected with pHEN phagemid (pHEN1-I or II), and plated on YTE, 100µg/ml ampicillin plates. Colonies were shaken at 37°C in 2xTY medium, 100 µg/ml ampicillin, 1% glucose to Ob₅₅₀=0.5 to 1.0. Cells were pelleted, washed once in 2xTY medium, resuspended in medium with 100 µg/ml ampicillin, 1 mM isopropyl β-D-thlogalactoside (IPTG), and grown for a further 16 hours. Cells were pelleted and the supernatant, containing the

secreted chains, used directly in ELISA.

The phagemid pHENI has the advantage over phage fd-CAT2, in that antibody can be produced either for phage 25 display (by growth in supE strains of E.coli) or as a tagged soluble fragment (by growth in non-suppressor strains), as a peptide tag (example 24) and amber codon were introduced between the antibody and g3p. Secretion of soluble Fab fragments from pHEN1-II or scFv fragments 30 from pHEN1-I was demonstrated after growth in E.coli HB2151 and induction with IPTG using Western blots (Figure 29). For detection of secreted proteins, 10ul supernatant of induced cultures were subjected to SDS-PAGE and proteins transferred by electroblotting to 35 Immobilon-P (Millipore). Soluble heavy and light chain were detected with goat polyclonal anti-human Fab antiserum (Sigma) and peroxidase conjugated rabbit antigoat immunoglobulin (Sigma), each at a dilution of The tagged VK domain was detected with 9E10 1:1000. antibody (1:1000) and peroxidase conjugated goat anti-40 mouse immunoglobulin (Fc specific) (1:1000) (Sigma) or with a peroxidase labelled anti-human CK antiserum (Dako). 3,3'-diaminobenzidine (DAB; Sigma) was used as peroxidase substrate (Harlow E., et al. 1988 Supr). With the scFv, the fragments were detected using the 9E10 45 anti-myc tag antibody (data not shown). With the Fab, only the light chain was detected by 9E10 (or anti-human CK) antibody, as expected, while the anti-human Fab antiserum detected both heavy and light chains. Binding 50 of the soluble scFv and Fab fragments to phOx-BSA (but not to BSA) was also demonstrated by ELISA (Table 5B).

Thus scFv and Fab fragments can be displayed on phage or

secreted as soluble fragments from the same phagemid vector.

Example 28 Increased Sensitivity in ELISA assay of Lysozyme using FDTscFvDl.3 as Primary Antibody Compared to Soluble scFvD1.3

In principle the use of phage antibodies should allow more sensitive immunoassays to be performed than with soluble antibodies. Phage antibodies combine the ability to bind a specific antigen with the potential for amplification through the presence of multiple (ca.2800) copies of the major coat protein (g8p) on each virion. This would allow the attachment of several antibody molecules directed against M13 to each virion followed by the attachment of several molecules of peroxidase-

15 conjugated anti-species antibody (anti-sheep) IgG in the Thus for every phage antibody bound to case below). antigen there is the potential for attaching several peroxidase molecules whereas when a soluble antibody is used as the primary antibody this amplification will not 20

occur.

ELISA plates were coated overnight at room temperature using 200µl of 10 fold dilutions of hen egg lysozyme (1000, 100, 10, 1, 0.1 and 0.01 µg/ml) in 50mM NaHCO3, pH9.6. ELISA was performed as described in 25 example 4 except that (i) incubation with anti-lysozyme antibody was with either FDTscFvD1.3 (pAb; 1011 phage per well; 1.6mol) or soluble affinity purified scFvDl.3 (18µg per well; 0.7nmol) (ii) incubation with second antibody was with 1/100 dilution of sheep anti-M13 serum for 30 FDTscFvD1.3 samples or with or 4/100 dilution of rabbit anti-scFvD1.3 serum (from S. Ward) for soluble scFvD1.3 samples (iii) peroxidase-conjugated rabbit anti-goat immunoglobulin (Sigma: 1/5000) was used for FDTscFvD1.3 samples and peroxidase-conjugated goat anti-rabbit 35 immunoglobulin (Sigma: 1/5000) was used for soluble scFvD1.3 samples. Absorbance at 405nm was measured after The results are shown in Figures 30 and 31. In these figures lysozyme concentrations for coating are shown on a log scale of dilutions relative to lug/ml. 40

(i.e. $\log = -3 = \log/ml$; $\log = 2 = 0.01 \, \mu g/ml$) Higher signals were obtained with FDTscFvDl.3 at all concentrations of lysozyme (Fig.31) but the difference was very marked at the greatest dilutions, where antigen quantities are most limiting (Figs. 30 and 31). 45 suggests that phage antibodies may be particularly

valuable for sandwich type assays where the capture of small amounts of antigen by the primary antibody will generate an amplified signal when phage antibodies directed against a different epitope are used as the

second antigen binding antibody. Example 29

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Direct Rescue and Expression of Mouse Monoclonal Antibodies as Single Chain Fv Fragments on the Surface of Bacteriophage fd.

The principle is very similar to that described in example 14. It consists of the PCR assembly of single chain antibodies from cDNA prepared from mouse monoclonals. As an example, the rescue and expression of two such antibodies from monoclonals expressing antibodies against the steroid hormone oestriol is described.

A. RNA Preparation

RNA can be prepared using many procedures well known to those skilled in the art. In this example, the use of Triton X-100 lysis, phenol/SDS RNase inactivation gave excellent results.

15 excellent results.

1. The mouse monoclonal cells that were used here had been harvested by centrifugation and resuspended in serum free medium. They were then centrifuged and resuspended

in saline and after a final centrifugation step, 20 resuspended in sterile water at 1 x 10° cells per ml. (Normally cells would be washed in PBS buffer and finally resuspended in PBS buffer, but these particular cells were supplied to us as described frozen in water.).

2. To 750µl of cells was added 250ul of ice cold 4X lysis buffer (40mM Tris HCl pH 7.4/4mM MgCl₂/600mM NaCl/40mM VRC (veronyl ribosyl complex)/2% Triton X-100).

The suspension was mixed well and left on ice for 5 minutes.

 Centrifugation was carried out at 4°C in a microfuge at 13000 rpm for 5 min.

The supernatant is then phenol extracted three times, phenol chloroform extracted three times and finally, ethanol precipitated as described in the materials and methods. The precipitate was resuspended in 50ul water.

35 4. The optical density of the RNA at 250nm with a 2.5ul sample in lml water was measured. The RNA was checked by electrophoresis of a 2ug sample on a 1% agarose gel. RNA in the range of 32ug to 42ug was obtained by this method. B. CDNA Preparation

The method used is the same as that described in example 14. Two cDNA preparations were made. These were from RNA extracted from the monoclonals known as cell lines 013 and 014 which both express antibodies against eh steroid hormone, cestriol.

45 C. Primary PCRS

C. Primary PCRs
The method used is essentially the same as that described in example 14. The VH region was amplified with the primers VHLBACK and VH1FOR-2. For the Vkappa region, four separate reactions were carried out using the primer VKZBACK and wither MJKIFONX, MJKZFONX,

50 the primer VK2BACK and wither MJK1FONX, MJK2FONX, MJK4FONX or MJK5FONX. Samples (5ul) were checked on a 1.5% agarose gel. From this it was observed that for

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cDNA prepared from the two oestriol monoclonals the primers VK2BACK and MJK1FONX gave the best amplification of the Vkappa region. The VH bands and the Vkappa bands amplified with VK2BACK/MJK1FONX were purified on 2% low melting point agarose gels for each monoclonals. The DNA bands were excised from the gel and purified using a dedicated Geneclean kit as described in example 14.

D. Preparation of linker
The method used is essentially the same as that
described in example 14. In this case, the amplified
linker DNA was purified on a 2% agarose gel and recovered 10 from the gel with a dedicated "Mermaid" kit (BIO 101, Geneclean, La Jolla, San Diego, California, USA) using the manufacturers instructions.

Assembly PCRs 15

The method used is essentially the same as that described in example 14. In this case, the assembled PCR product was purified on a 2% agarose gel and recovered from the gel with a dedicated "Mermaid" kit.

Adding restriction sites and work-up The assembled product was "tagged" with Apa LI and Not I restriction sites. The DNA was then digested with Apa LI and Not I to give the appropriate sticky ends for cloning and then purified on a 2% low melting point agarose gel and extracted using a Geneclean kit. The method used is the same as that described in example 14.

Cloning into Vector fd-CAT2

A total of 15ug of CsCl purified fd-CAT2 DNA was digested with 100 units of the restriction enzyme Not I (New England Biolabs) in a total volume of 200ul 1X NEB Not I buffer with 1% NEB acetylated BSA for a total of 3 hours at 37°C. The vector DNA was the treated twice with 15ul Strataclean (a commercially available resin for the removal of protein), following the manufacturers instructions (Stratagene, 11099 North Torrey Pines Road, The DNA was then ethanol La Jolla, California, USA). precipitated and redissolved in TE buffer (Sambrook et al., 1989 supra). The DNA was then digested with 100 units of the restriction enzyme Apa LI (New England Biolabs) in a total volume of 200ul 1X NEB Buffer 4

40 overnight at 37°C. The vector was then purified with a Chroma Spin 1000 column following the manufacturers instructions (Clontech Laboratories Inc. 4030 Fabian way, Palo Alto, California, USA). This step removes the Apa LI/Not I fragment to give cut vector DNA for maximum 45

ligation efficiency.

Ligation reactions were carried out with 2.5-10ng of the DNA insert and long of vector in a total volume of 10ul of 1% NEB ligase buffer with lul of NEB ligase (New England Biolabs) at 16°C overnight (approx 16 hours). Transformation and growth

E.coli strain TG1 was made competent and transformed

with the fdCAT2 recombinant DNA as described by Sambrook et al, 1989 Supra. The cells were plated out on LBtet plates (10g tryptone, 5g yeast extract, 10g NaCl, 15g bacto-agar per litre with 15ug/ul of tetracycline added just before pouring the plates) and grown overnight.

Single well isolated colonies were then inoculated into 10 ml of LBtet broth (LB medium with 15ug/ul of tetracycline) in 50 ml tubes. After overnight growth at 35°C/350rpm in a bench top centrifuge. The supernatants were transferred to 15 ml centrifuge tubes and 2ml 20% PEG 8000/2.5M NaCl added to each. After incubating at room temperature for 20-30 minutes, the recombinant phage was pelleted by centrifugation at 9000rpm in a Sorval SM24 rotor for 30 minutes. The PEG supernatant was

15 discarded. Any remaining PEG was removed with a pasteur pepette after a brief (2 minutes) centrifugation sten. This last step was repeated to make sure that no PEG remained. The phage pellet was then resuspended in 500ul PBS buffer. This was transferred to a microcentrifuge 20 tube and spun at 13000 rpm to remove any remaining cells.

The phage supernatant was transferred to a fresh tube.

Assay for antibody expression

Bacteriophage fd recombinants were screened for the expression of antibody against cestriol by ELISA. 25 method is described in example 6. In this case the following alterations are relevant.

Microtitre plates were coated overnight with 40ug/ml oestriol-6 carboxymethyloxime-BSA (Staraloids. 31

Radcliffs Road, Croydon, CRO 5QJ, England).

1st antibody was the putative phage anti oestriol antibody. 50ul of phage in a final volume of 200ul of 30 sterile PBS combining 0.25% gelatin was added to each well.

3. 2nd antibody was sheep anti M13 at 1:1000 dilution. 35 4. 3rd antibody was peroxidase conjugated rabbit anti

goat immunoglobulin.

Recombinants expressing functional antibody were detected by incubation with the chromogenic substrate 2'2' axinobis (3-ethyl benzthiazoline sulphonic acid). The results are shown in figures 32 and 33.

40 Example 30

Kinetic Properties of Alkaline Phosphatase Displayed on the Surface of Bacteriophage fd

This example demonstrates that kinetic properties of 45 an enzyme expressed on phage are qualitatively similar to those in solution. Bacteriophage fd displaying alkaline phosphatase fusions of gene 3 with either the native arginine (see example 31) or the mutant residue alanine at position 166 (see example 11) were prepared by PEG 50 precipitation as described in the materials and methods.

The kinetic parameters of alkaline phosphatase expressed on the surface of fd phage were investigated in

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1M Tris/HCl, pH8.0 at 20°C with 1ml 4-nitrophenyl phosphate as substrate. The reactions were initiated by the addition of 100µl of a phage-alkaline phosphatase fusion preparation, 50 fold concentrated with respect to the original culture supernatant. The rate of change of absorbance was monitored at 410nm using a Philips 8730 spectrophotometer and the initial reaction rate calculated using a molar absorbance of 16200 1/mol/cm. For the fdphoAla 166 enzyme but not fdphoArg166 a lag phage was seen following this addition, the reaction rate accelerating until a steady state was obtained after approximately 60 to 90 secs. This steady state rate was used for determination of kinetic parameters. deviation form Michaelis Menten kinetics was apparent for either phage enzyme. Values of K_{m} and k_{cat} were derived from plots of s/v against s and are shown in Table 6.

Because of the difficulty in establishing the relationship between the number of phage particles an the number of active enzyme dimers formed on the phage koat values are expressed not as absolute values, but as relative values between the two enzyme forms. Western blots (carried out as in example 31 using antig3p antiserum) of the phage enzyme preparations used in this experiment showed approximately equal intensities for the full length fusion band with the Arg166 and Ala166 enzymes when detected using antibody directed against In these preparations the intact fusion represents approximately 30% of the detected material. The two preparations were therefore assumed to be expressing approximately the same concentrations of

intact fusions. Table 6 summarises the kinetic data from this experiment and compares it with data from Chaidaroglou, A. et al (Biochemistry 27, 8338-8343 (1988)) obtained with soluble preparations of the wild type and mutant enzyme forms. The same substrate and assay conditions were used in both experiments. Soluble alkaline phosphatase was also tested in parallel in our experiments (Km=8.5µM; kcat=3480 mol substrate converted mol enzyme 1 min 1).

The effect of mutating arginine at position 166 to alanine is qualitatively similar for the phage enzyme as for the soluble enzyme. Km is increased about 15 fold and the relative kcat is decreased to 36% of that for wild type. This increased $K_{\rm IM}$ would reflect a reduction in substrate affinity in the phage enzyme on mutation of Arg166, as was proposed for the soluble enzyme (Chaidaroglou et al, 1988 supra), assuming the same kinetic mechanism applies. There are, however, some quantitative differences in the behaviour of K_{Γ} of the 50 phage enzyme. The $\rm K_m$ of 73µM observed for fdphoArg166 compares with a $\rm K_m$ of 12.7µM for the free enzyme; the $\rm K_m$

for fdphoAlal66 is 1070µM whereas the free mutant enzyme has a $K_{\rm m}$ of 1620µM. One can speculate that the higher $K_{\rm m}$ for fdphoArg 166 and the lower $K_{\rm m}$ for fdphoAlal66, compared to the soluble enzymes result from the 'anchored' alkaline phosphatase fusion molecules interacting to form dimers in a different manner to the enzyme in free solution.

The relative values of k_{Cat} for the Arg166 and Ala166 forms are however very similar for both the phage of enzymes and the soluble enzymes, a reduction occurring on mutation to 35 to 40% of the value for the native enzyme. The rate limiting step, determining k_{Cat} for soluble phoArg166 is thought to be dissociation of non-covalently

bound phosphate from the enzyme (Hull W.E. et al. Biochemistry 15, 1547-1561 1976). Chaidaroglou et al (1988) supra suggest that, for the soluble enzyme, mutation of Argl66 to alenine alters additional steps, one of which may be hydrolysis of the phosphoenzyme intermediate. The similarity in the reduction in k_{Cat} on mutation of Argl66 to alenine for the phage enzymes suggests that the same steps may be altered in a

suggests that the same steps may be altered in a quantitatively similar manner in the mutant phage enzyme as in the mutant soluble enzyme.

Thus, enzymes displayed on phage show qualitatively similar characteristics to soluble enzymes.

Example 31

Demonstration using Ultrafiltration that Cloned Alkaline Phosphatase Behaves as Part of the Virus Particle

The construct fdphoAlal66 (derived in example 11)
was converted back to the wild type residue (arginine) at
position 166 by in vitro mutagenesis (Amersham
International) using the printer

APARG166:5' TAGCATTTGCGCGAGGTCACA 3'.

This construct with the wild type insert was called 35 fdphoArgl66.

E.coli TGl or KS272 cells (cells with a deletion in the endogenous phoA gene, Strauch and Beckwith, 1988 Supra) containing either fd-phoAlal66, fdphoArgl66 or fd-CAT2 were grown for 16 hours at 37°C in 2xTY with 15µg/ml tetracycline. Concentrated phage were prepared as follows. Phage-enzyme cultures are clarified by centrifugation (15 min at 10,000 rpm, 8 x 50 ml rotor, sorval RC-58 centrifuge). Phage are precipitated by

adding 1/5 volume 20% polyethylene glycol, 2.5 M Nacl, 1eaving for 1 hr at 4°C, and centrifuging (as above). Phage pellets are resuspended in 10 mM Tris-HCl, pH 8.0 to 1/100th of the original volume, and residual bacteria and aggregated phage removed by centrifugation for 10 to 15 minutes in a bench microcentrifuga at 13000 rpm at

50 4°C.

SDS/Polyacrylamide gel electrophoresis and western blotting were basically as described previously (example

Denatured samples consisting of 16µl of a 50 fold

concentrate of phage were separated using a 10% SDS/polyacrylamide gel and detected with polyclonal antiserum raised against either E.coli alkaline phosphatase (Northumbria Biologicals, South Nelson Industrial Estate, Cramlington, Northumberland, NE23 9HL) or against the minor coat protein encoded by gene 3 (from Prof. I. Rasched, Universitat Konstanz, see Stengele et al, 1990) at 1 in 1000 dilution. This was followed by

al, 1990) at 1 in 1000 dilution. This was followed by incubation with peroxidase-conjugated goat-anti-rabbit immunoglobulin (Sigma I in 5000) and detection with the ECL Western blotting system (Amersham International).

The presence of fusion proteins was confirmed by

western blotting of proteins from phage particles derived from fd-phoklai66 (phage-enzyme) or fd-CAT2 (vector phage). Detection with antiserum raised against the gene 3 protein reveals a product of apparent relative molecular mass (Mr) of 63,000 in vector phage (figure 34e). Although this is different from the predicted

molecular weight based on the amino acid sequence (42,000), the natural product of gene 3 has previously been reported to exhibit reduced mobility during

electrophoresis (Stengele et al, 1990).

In the fd-phoAlal66 sample the largest band has an apparent Mr of 115,000, (fig. 34). Taking into account the aberrant mobility of the gene 3 portion of the fusion, this is approximately the size expected from fusing with an alkaline phosphatase domain of 47 kD. This analysis also reveals that a proportion of the Gene3

30 reactive material in this phage-enzyme preparation is present at the size of the native gene3 product, suggesting that degradation is occurring. In the preparation shown in figure 34, approximately 5-10% of the gene 3 fusions are intact. In more recent

35 preparations and in all the preparations used in this example and example 32, approximately 30-60% of fusions

are full length.

The protein of Mr 115,000 is the major protein observed in Western blots of phage-enzyme derived from 40 TG1 cells when probed with antiserum raised against E.coli alkaline phosphatase (anti-BAP), confirming the assignment of this band to intact fusion. Further, when phage enzyme is prepared using KSZ72 cells, which have a deletion in the endogenous phom gene (Strauch & Beckwith,

45 1988, supra.) it is also the major band. There are additional bands at Mr 95000 and 60000 reactive with anti-BAP antiserum which may indicate degradation of the

fusion product.

The anti-BAP antiserum also reacts wit material running with the dye front and with a molecule of Mr 45,000 but evidence suggests that this material is not alkaline phosphatase. This pattern is detected in PEG

precipitated vector phage samples (figure 34c) and is not therefore contributed by protein expressed from the cloned phoA gene. These bands are detected in culture supernatants of cells carrying fd-CAT2 but is not detected in the supernatant of uninfected cells (not shown) and so either represents cross-reactivity with phage encoded material or with a PEG precipitable cellular component leaked from infected cells (Boeke et al. Mol. Gen. Genet. 186, 185-192 1982). Although the 10 fragment of Mr, 45,000 is close to the size of free alkaline phosphatase (47,000), it is present in phage preparations from KS272 cells which have a deletion in Furthermore its mobility is different the phoA locus. from purified alkaline phosphatase and they can be 15 distinguished by electrophoresis (figure 34d).

Ultrafiltration was used to confirm that the fusion protein behaved as though it were part of a larger structure, as would be expected for an enzyme bound to a phage particle. Phage samples (100µl of a 50 fold concentrate) were passed through ultrafiltration filters with a nominal modecular weight limit of 300000 deltons (Ultrafree-MC filters, Millipore) by centrifugation for 5 to 15 minutes at 13,000 r.p.m. in an MSE microcentaur microfuge. Retained material was recovered by

25 resuspending in 100µl of 10mM Tris, pH 8.0.

Phage-enzyme or free alkaline phosphatase (83ng) mixed with vector phage were passed through filters with a nominal molecular weight limit of 300,000 daltons (Ultrafree-MC filters, Millipore). Figure 35 A again shows that the band of Mr. 115,000 is the major product 30 reactive with anti-BAP antiserum. This and the other minor products reactive with anti-BAP are present in material retained by the ultrafiltration membrane. Analysis of retained and flow through fractions of phage 35 preparations derived from KS272 demonstrates that different molecular species are being separated by the ultrafiltration membranes. Figure 35b shows the protein of Mr 115,000 is retained by the filter whereas the putative degradation products of Mr 95,000 and 60,000 found in phage preparations derived from KS272 cells, are 40 not retained.

In mixture of alkaline phosphatase and vector phage Figure 35c-f, free alkaline phosphatase (dimer size of 94,000 daltons) is detected in the flow through as a monomer band with Mr 47,000 on denaturing polyacrulamide gels (figure 35B), while the cross reactive molecule found in vector phage preparations (Mr 45,000) is in retained on the filter (figure 35B). This suggests that the cross reactive molecule is part of the phage particle and underlines the fact that the ultrafiltration membranes are effecting a separation. Thus the expected fusion band in this phage-enzyme is present in material

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retained on ultrafiltration membranes demonstrating that it is part of a larger structure as would be expected for

viral bound enzyme.

Catalytic activity has been demonstrated on phage particles expressing alkaline phosphatase. Table 7 shows that the wild type alkaline phosphatase gene expressed on phage (fd-phoArgl66) has a specific activity (moles of substrate converted per mole of viral particles) of This is close to the turnover value of 3.700/min. 4540/min found for purified alkaline phosphatase by 10 Malamy and Horecker, Biochemistry 3, 1893-1897 1964).

Chaidaroglou et al, 1988 supra have shown that substituting alanine for arginine at the active site (residue 166) leads to a reduction in the rate of Preparations of phage displaying alkaline catalysis. phosphatase with this mutation derived from TG1 and KS272 show reduced specific activities of 380 and 1400 mol substrate converted/mol phage/min respectively. Enzyme activity was measured in the retained and flow-through fractions prepared by ultrafiltration, shown in figure The bulk of activity from phage-enzyme was retained on the filters whereas the majority of activity from free enzyme passes through. Therefore, the enzyme activity in these fusions behaved as would be expected for virally associated enzyme (not shown). Little or no catalytic 25 activity is measured in preparations of vector phage from either TGI or KS272 cells (Table 7), indication that the catalytic activities above are due to phage enzyme and not contamination with bacterial phosphatase. of phage particles to soluble enzyme does not have a 30

significant effect on activity (Table 7). Therefore, both the catalytic and immunochemical activity of alkaline phosphatase have been demonstrated to be due to enzyme which is part of the phage particle.

Example 32

Affinity chromatography of phage alkaline phosphatase Affinity chromatography, using the specific binding properties of enzymes has proved to be a very powerful The purification of method for their purification. phage-enzymes by this approach would enable the genetic material encoding the enzyme to be isolated with the enzyme itself. Thus, mutagenesis of cloned enzymes expressed on the surface of filamentous bacteriophage will lead to a whole population of enzyme variants, from

which variants with desired binding properties could be 45 isolated. Soluble alkaline phosphatase (from calf intestine)

has been purified by binding to immobilised arsenate (a competitive inhibitor), and eluting with inorganic phosphate, which is a product (and competitive inhibitor) of the enzyme reaction (Brenna, O. et al, Biochem, J. 151 291-296 1975). The applicants have determined that

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soluble alkaline phosphatase from E.coli is also retained by this matrix (not shown). In this example it is demonstrated that phage displaying E.coli alkaline phosphatase binds to arsenate-Sepharose and can be specifically eluted.

Arsenate-Sepharose was prepared by coupling 4-(paminophenylazo) phenyl arsonic acid to tyraminyl-Sepharose according to the method of Breena et al. (1975; supra). Affinity chromatography of phage enzyme fdphoArgl66 (example 31) was carried out in a disposable chromatography column with a 0.5 ml column volume. Columns were prewashed with 100 volumes of column buffer (100mM Tris pH 8.4, 1mM MgCl2, 0.1 mM ZnCl2, 0.1% Tween 20, Brenna et al, 1975, supra.) lml of a 40 fold concentrate of phage-enzyme (in column buffer; prepared as in example 31) was loaded and washed through with 100 volumes of column buffer. Bound phage-enzyme was eluted with 5mls of column buffer containing 20mM NaHPO4. eluate and wash fractions were quantitated by dot blotting onto nitrocellulose and comparing with known amounts of phage-enzyme. The blots were detected using sheep anti-M13 antiserum (gift from M. Hobart), antisheep peroxidase (Sigma) and enhanced chemiluminescent substrate (Amersham). A range of exposures were taken.

25 Table 8 shows the results of affinity chromatography of phage displaying alkaline phosphatase on arsenate-Sepharose. In separate experiments phage particles expressing either mutant (fdphoAla 166; example 11) and or wild type (fdphoArg 166) forms are retained on 30 arsenate-Sepharose and eluted with inorganic phosphate. Approximately 0.5 to 3% of added phage enzyme particles loaded ('input phage') were specifically eluted with phosphate ('output phage') compared to only 0.05% of vector particles. Arsenate is a competitive inhibitor 35 with K; of 20µM with respect to 4- nitrophenyl phosphate. Phage particles antibodies have previously been isolated on the basis of interactions with similar affinities (example 23). This association is in within the range of a large number of enzyme-ligand interactions suggesting 40 wide applicability for this approach.

Table 8 also shows that the infectivity of phage particles expressing enzyme is reduced with compared with vector phage particles. This makes titration of infectious particles an inappropriate means of quantitating the number of phage enzyme particles. For this reason the number of phage were measured by dot blotting and phage were detected with anti-M13 antiserum as above.

Whereas, overall recovery of catalytic activity may be an important consideration in enzyme purification, this is not critical with phage-enzymes. Even if only low levels of phage-enzyme bind to and are specifically

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eluted from affinity columns, this will generate clones which can subsequently be grown up in bulk as phage-enzymes or can be transferred to expression vectors yielding soluble products.

Example 33
PCR Assembly of DNA encoding Fab Fragments of an Antibody

directed against Oxazolone

Example 25 showed that genes encoding Fab fragments could be subcloned into vectors fdCAT2 and pHEN1 and the protein domains displayed on the surface of phage with retention of binding function. This example shows that the VMCH and VKCK domains can be amplified separately and then joined by a linker allowing the expression of the light chain as a geneIII protein fusion and the VMCH.

fragment as a soluble molecule. A functional Fab fragment is then displayed on phage by association of these domains. The assembly process, described in this example, is required for display of a library of Fab fragments derived from the immune repertoire if both heavy and light chain domains are to be encoded within a

single vector.

The VHCH1 and VKCK domains of a construct (example 25; construct II in pUC19) derived from antibody NQ10 12.5 directed against 2-phenyl-5-oxazolone were amplified using PCR. For cloning into the vector fdCAT2 the cligonucleotides VHIBACKAPA (example 25) and HuIgG1-4 CHIFOR (example 40) were used to amplify the VHCH1 domains. For cloning into pHEN1 VHIBACKSFH5 (example 25)

replaced VHIBACKAPA for this amplification. For cloning into both vectors the VKCK domains were amplified using VKZBACK (example 25) and CKNOTFOR (example 40). A linker oligonucleotide fragment containing the bacteriophage fd gene 8 terminator and the fd gene 3 promoter was prepared by amplifying the region containing them from the vector

35 fdCAT2 by PCR using the oligonuclectides.

VK-TERM-FOR

5' TGG AGA CTG GGT GAG CTC AAT GTC GGA GTG AGA ATA GAA AGG 3' (overlapping with VK2BACK [example 14]) and

40 CHI-TERM-BACK
5'AAG CCC AGC AAC ACC CTG GAC AAG AAA GTT GAG CCC AAA
TCT AGC TGA TAA ACC GAT ACA ATT AAA GGC 3' (overlapping

with HulgG1-4 CH1-FOR)

Assembly of the Fab fragment from the amplified VHCH1 and 45 VKCK domains and the linker prepared as above was as described in example 14E except that the primers VHIBACKAPA (when cloning into fdCAT1) or VHIBACKSFH5 (when cloning into pHEN1) and CKNOTFOR were used for the final reamplification, thereby introducing restriction

sites for cloning into fdCAT2 (Apall-NotI) or pHEN1 (Sfil-NotI) the assembled Fab fragment is shown in figure 34. No assembled product was seen in the absence of linker. An assembled scFv prepared according to example 14 is shown for comparison.

Phage antibodies were prepared as in example 25 and ELISA was performed with oxazolone as antigen according to example 6. Results were as expected for Fab fragments cloned in both fdCAT2 and pHEN1 samples, phage particles bound to oxazolone as detected by a positive ELISA signal.

Example 34

10 Construction of a Gene III Deficient Helper Phage

To fully realise the potential of the phagemid cloning system, a helper phage lacking gene III is desirable. Rescue of gene III fusions with such a helper phage would result in all the progeny phagemids having a gene III fusion on their capsid, since there would be no competition with the wild type molecule.

Control over the number of fusion molecules contained on each phage will provide particularly useful. For example, a gene III deficient helper phage can be

20 used to rescue low affinity antibodies from a naive repertoire, in which high avidity will be necessary to isolate those phage bearing the correct antibody specificity. The unmutated helper phage can then be used when higher affinity versions are constructed, thereby reducing the avidity component, and permitting selection purely on the basis of affinity. This will prove a surprisingly successful strategy for isolation and affinity maturation of antibodies from naive libraries.

The strategy chosen to construct the helper phage was to partially delete gene III of MI3KO7 using exonuclease Bal 31. However, phage lacking gene III protein are non-infective so an E.coli strain expressing gene III was constructed. Wild type MI3 gene III was FCR-amplified with primers gIIIFUFO and gIIIFUBA, exactly

35 as described in exemple 24. The FCR product was digested with Eco RI and Hind III and inserted into Eco RI and Hind III and inserted into Eco RI and Hind III. The FCR product was digested with Eco RI and Hind III and inserted into Eco RI and filamentous phage origin of SS DNA replication) under control of the lac promoter. The plasmid was transformed into E.coli TG1, and the resulting strain called TG1/pUCl9gIII. This strain provides gIII protein in

trans to the helper phage.

There is a single unique Bam HI site in M13KO7.

which is approximatively in the centre of gIII. Doublestranded M13KO7 DNA was prepared by alkaline lysis and caesium chloride centrifugation (Sambrook et al. et supra. 1989); twenty up of DNA was cut with Bam H1, phenol extracted and ethanol precipitated then resuspended in 50µl of Bal 31 huffer (600mM NaCl, 20mM FDM). Tris-HCl pH 8.0. 12 mM CaCl. 12mm MCCl and hww FDM).

50 Tris-HCl pH 8.0, 12 mM CaCl₂, 12mM MgCl₂ and 1mM EDTA) and digested for 4 minutes with 1 unit of Bal 31 (New England BioLabs). This treatment removed approximatley

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IKb of DNA. EGTA was added to 20mM and the reaction phenol extracted and ethanol precipitated prior to purification of the truncated genome on an agarose gel. The DNA was repaired with Klenow enzyme and self-ligated with T4 DNA ligase (New England BioLabs).

Aliquots of the ligation reaction were transformed into competent TGI/pUCl9gIII and plated on SOB medium containing ampicillin at 100µg/ml and kanamycin at 50µg/ml. Colonies were screened for the presence of a deletion by PCR with primers gIIIFUBA and KSJ12

10 deletion by PCR with (CGGAATACCCAAAGAACTGG).

KSJ 12 anneals to gene VI which is immediately downstream of gIII in the phage genome, so distinguishing gIII on the helper phage from that resident on the plasmid. Three clones gave tructated PCR products corresponding to deletions of ca. 200, 400 and 800bp. These clones were called M13KO7 gIII \(\Lambda \) Nos 1,2 and 3 respectively. No clones were isolated from the earlier Bal 31 time points, suggesting that these are in some way lethal to the host cell. Several clones were isolated from later time points, but none of these gave a PCR product, indicating that the deletion reaction had gone

too far.

MISKO7 gIII \(\text{No.s} \) 1,2 and 3 were cultured and the resulting helper phage tested for their ability to rescue an antibody gIII fusion (scFv Dl.3) by ELISA, exactly as described in example 18. As shown in figure 37, only one clone, MISKO7 gIII\(\text{NO3} \) NO3 was found to rescue the antibody well; in fact the signal using this helper was greater than that observed with the parent MI3 KO7. MI3KO7 gIII\(\text{NO3} \)

than that observed with the parent M13 KO7. M13KO7 gill?
No3 rescued phagemids should have a much higher density
of antibody fusions on their surfaces. That this was
indeed the case was demonstrated when the phage used in
this ELISA were analysed by Western blotting with anti
gill protein antiserum (fig. 38). This analysis enables

35 gIII protein antiserum (fig. 38). This analysis enables estimation of the amount of gIII fusion protein versus free gIII protein present on the phage(mid) particles. Only a minute fraction of the gIII protein on the

M13K07-rescued material is present as an intact fusion (fig 38). The fusion protein band is induced by IPTG, so is indisputably that synthesized by the phagemid. As expected, even when the lac promoter driving gIII fusion protein synthesis is fully induced (100µM IPTG), wild type gIII protein, at a lower copy number and driven from

a far weaker promoter, predominates. This is in contrast to the pattern generated by the same clone rescued with M13KO7 gIII/NO3, and the pattern generated by fd CAP2-scFv Dl.3. In both of these latter cases, there is no competition with wild-type gIII and the fusion protein band is correspondingly stronger.

It is worthy of note that construction of M13K07 qIII A No3 was immensely inefficient: one clone from 20µg

of starting DNA. Moreover, the yield of gIII helper phage from overnight cultures is extremely low ca.10⁵ cfu/ml compared with ca. 10¹¹ cfu/ml for the parental phage. Despite this, MI3RO7 gIII NO3 rescues the phagemid as well as the parental phage, as judged by the number of phagemid particles produced after overnight growth. This indicates that trans replication and packaging functions of the helper are intact and suggest that its own replication is defective. Hence it may be that inactivation of gIII is normally toxic to the host cell, and that MI3RO7 gIII A NO3 was isolated because of a compensating mutation affecting, for example, replication. Phage fd-tet is unusual in that if

replication. Phage fd-tet is unusual in that it tolerates mutations in structural genes that are normally 15 lethal to the host cell, since it has a replication defect that slows down accumulation of toxic phage products; MISKOT gIII/A No3 may also have such a defect.

Mi3KO7g IIIA No 3 has been deposited at the National Collection of Type Cultures, 61 Colindale Avenue, London, NW9 6HT, UK (Accession No. NCTC 12478). On 28 June 1991, in accordance with the regulations of the Budapest Treaty. It contains a deletion of the MM3 genome from bases 1979 to 2768 inclusive (see Van Wezenbeek, F.G.M.F. et al., Gene II p129-148, 1980 for the DNA sequence of

25 the M13 genome). Example 35

Selection of bacteriophage expressing scFv fragments directed against lysozyme from mixtures according to affinity using a panning procedure

For isolation of an antibody with a desired high affinity, it is necessary to be able to select an antibody with only a few fold higher affinity than the remainder of the population. This will be particularly important when an antibody with insufficient affinity

35 has been isolated, for example, from a repertoire derived from an immunised animal, and random mutagenesis is used to prepare derivatives with potentially increased affinity. In this example, mixtures of phage expressing antibodies of different affinities directed against hen egg lysozyme were subjected to a panning procedure. It is demonstrated that phage antibodies give the ability to select for an antibody with a R₂ of 2nM against one with

a K_d of 13nM.

The oligonucleotides used in this example are shown in the list below:
OLIGONUCLEOTIDES

VHEHDI3APA : 5'- CAC AGT GCA CAG GTC CAA CTG CAG GAG AGC GGT
WHFHDI3 : 5'- CGG TGA CGA GGC TGC CTT GAC CCC

VHFHD13 : 5'- CGG TGA CGA GGC TGC CTT GAC CCC
HD13BLIN : 5'- GGG GTC AGG GCA GCC TCG TCA CCG
HD13FLIN3 : 5'- TGG GCT CTG GGT CTT GGAT GTC CGA T
VKBHD13 : 5'- GAC ATC CAG ATG ACC CAG AGC CCA

VKFHD13NOT : 5'- GAG TCA TTC TGC GGC CGC ACG TTT GAT TTC CAC CTT GGT CCC MURD13SEQ : 5'- GAG GAG ATT TTC CCT GT

HUMDISSEQ : 5'- TTG GAG CCT TAC CTG GC

5 FDPCRFOR : 5'- TAG CCC CCT TAT TAG CGT TTG CCA FDPCRBAK : 5'- GCG ATG GGT GTT GTC ATT GTC GGC

Phage displaying scFv fragments directed against lysozyme were derived from cloned Fab fragments in plasmids.

Heavy and light chain variable regions were amplified by the polymerase chain reaction (PCR) from plasmids containing humanized VH-CH1 or VK-CK inserts suitable for production of Fab fragments (gift of J. Foote). The dissociation constant, Kd for different combinations of the two plasmids combined as Fabs, are

 shown below:
 Light Chain Plasmid
 Kd

 Heavy Chain Plasmid
 Light Chain Plasmid
 Kd

 HuH-1
 HuK-3
 52 nM

 HuH-1
 HuK-4
 180 nM

 HuH-2
 HuK-3
 13 nM

 HuH-2
 HuK-4
 (not determined)

Primary PCR of the variable regions was performed by combining the following:

5 36.5 µl Water

5 µl PCR buffer (10x)

2 µl dNTP (5mM)
2.5 µl Back oligo (10 pmoles/µl) (VHBHD13APA or VKBHD13)
2.5 µl Forward oligo (10 pmoles/µl) (VHFHD13 or VKFHD13NOT)

VKFHDI3NOT)

The reaction is decontaminated by UV irradiation to destroy foreign DNA for 5 minutes, and 1 µl of plasmid DNA added (0.1 µg/µl). The por mixture was covered with 2 drops of paraffin oil, and placed on the por block at 35 94°C for 5 minutes before the addition of 0.5 µl of Tag DNA polymerase under the paraffin. The cycling conditions used were 94°C 1 min, 40°C 1 min, 72°C 1.5 min 17 cycles.

The linker (Gly4-Ser)3, was amplified from the antiphox (2-phenyloxazol-5-one) clone fd-CAT2-SeFv NQ11,
using the oligos HD13BLIN and HD13FLIN3, with 0.lpg of
plasmid DNA. The PCR cycling used was 94°C 1 min, 23°C
1.5 min, for 17 cycles.

Amplified DNA was purified by running the samples on 45 a 2% low melting point agarose gel at 90 mA, excising the appropriate bands and extracting the DNA using the Geneclean II kit (BIO 101 Inc.) for the VH and VK, or by using Spin-X filter units (Costar) for the linker. A final volume of 10 µl was used to resuspend the extracted

50 DNA. PCR Assembly

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Assembly of the four single chain Fv Humanized D1.3

(scFv HuD1.3) constructs was by the process of 'assembly by overlap extension' example 14.

The following were combined: 34.5 ul Water

5 µl PCR Buffer (10x)

2 µl dNTP (5 mM)

2.5 µl Back oligo (10 pmoles/ul) (VHBHD13APA) 2.5 ul Forward oligo (10 pmoles/ul) (VKFHD13NOT)

Once again, the reaction is decontaminate by UV 10 treatment for 5 minutes before the addition of 1 µl of the primary PCR products; VH-1 or VH-2, VK-3 or VK-4, plus the linker DNA. The reaction was covered with 2 drops of paraffin, and heated at 94°C for 5 minutes before the addition of 0.5 µl of Tag Polymerase. The PCR 15

cycling conditions used were 94°C 1 min. 60°C 1.5 min. 72°C 2.5 min for 20 cycles.

The aqueous layer under the paraffin was extracted once with phenol, once with phenol: chloroform, once with ether, ethanol precipitated, and resuspended in 36 µl of

20 water. To this was added, 5 µl of 10x Buffer for Not1, 5 µl 1 mg/ml BSA, and 4 µl (40 U) of Not1 (New England Biolabs). The restriction was incubated at 37°C overnight.

The DNA was ethanol precipitated and resuspended in 25 36 μ l of water, and 5 μ l 10x NEB Buffer 4, 5 μ l 1 mg/ml BSA, and 2 μ l (40 U) of ApaLI (New England Biolabs). This was incubated at 37°C for 5 hours; a further 2 ul of ApaLI was added and the reaction incubated at 37°C overnight.

The cut DNA was extracted by gel purification on a 1.3% low melting point agarose gel followed by treatment 30 with Geneclean, to yield the insert DNA for cloning.

Vector fd CAT2 (prepared and digested with ApaLI and NotI as in example 20) and the scFv DNA were ligated as 35 in example 20.

Analysis Of Clones

Colonies from the ligations were first screened for inserts by PCR screening. The PCR mixture was prepared in bulk by combining 14.8 µL 1x PCR Buffer, 1 µl dNTP (5 40 mM), 1 µl Back oligo (FDFCRBAK), 1 µl Forward oligo (FDPCRFOR), and 0.2 µl Tag polymerase per colony screened. 20 pl of this PCR mixture was aliquoted into a 96 well Techne plate. The top of a colony was touched with a toothpick and twirled quickly into the PCR mixture

45 and the colony rescued by placing the toothpick in a Cellwell plate (Nunc) containing 250 µl of 2x TY medium. The PCR mixture is covered with 1 drop of paraffin and the place placed on the block at 94°C for 10 minutes before cycling at 94°C 1 minute, 60°C 1 minute, 72°C 2.5 50

minutes.

The clones thus derived were named as below. affinity of scFv fragments derived the Fab fragments was not determined but previous results suggests that these are closely related although not necessarily identical (R.E. Bird & B.W. Walker TIBTECH 9 132-137, 1991).

5	Construct Name	Composition	Affinity of Fab (Kd)
	TPB1	VH-HuH2-(Gly4-Ser)3-VK-HuK3	13 nM
	TPB2	VH-HuH1-(Gly4-Ser)3-VK-HuK4	180 nM
10	TPB3	VH-HuH2-(Gly4-Ser)3-VK-HuK4	(Unknown)
	TPB4	VH-HuH1-(Gly4-Ser)3-VK-HuK3	52 nM

Preparation of phage and ELISA was as described in example 6. The clones generated in fd CAT2 were shown to bind lysozyme as expected. 15

Affinity selection Selection of Highest Affinity Binding Phage

Mixing experiments were performed in which fd-CAT2 scrvDl.3 phage (example 19) were mixed with either fd-CAT2 TPB1, fd-CAT2 TPB2, or fd-CAT2 TKPB4, and used in 20 one round of panning.

The general method used for affinity selection by panning is that detailed below. Any deviation from this protocol is described at the relevant point. Panning plates were placed on a rocking platform between

manipulations.

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Falcon 35 mm Tissue Culture dishes were coated overnight with 1 ml of Lysozyme (various concentrations) dissolved in 50 mM Sodium Hydrogen Carbonate, pH 9.6, and blocked with 2 ml 2% MPBS at room temperature for 2 30 hours. Phage were prepared in 1 ml 2% MPBS and rocked at room temperature for 2 hours. Plates were washed for 5 minutes with 2 ml of the following solutions; 5 times with PBS, PBS-Tween, 50 mM Tris-HCl, pH 7.5; 500 mM Sodium Chloride, 50 mM Tris-HCl, pH 8.5; 500 mM Sodium 35 Chloride, 50 mM Tris-HCl, pH 9.5; 500 mM Sodium Chloride, 50 mM Sodium Hydrogen Carbonated, pH 9.6; 500 mM Sodium Chloride. Phage were then eluted by adding 1 ml 100 mM Triethylamine and rocking for 5 minutes before removing the eluate which was neutralised with 100 ul 1.0 M Tris-40 HC1, pH 7.4.

Plates were coated overnight with Lysozyme at the

concentration listed below.

Colonies from the single round of panning were probed with either MURDSEQ (for fdCAT2 scFvD1.3) or 45 HUMDISSEQ (for fdCAT2 TPB constructs).

Circles of nitrocellulose (Schleicher & Schuell, BA 85, 0.45 µm) were labelled in pencil and lowered gently onto the colonies derived from the panning experiments and left for one minute. The filters were then pulled 50 off quickly from one edge and placed colony side up on a piece of 3MM paper (Whatman) soaked in Denaturing

solution (500 mM Sodium Hydroxide: 1.5 M Sodium Chloride) for 5 minutes. They were then transferred to 3MM soaked in Neutralizing Solution (3.0 M Sodium Chloride: 500 mM Tris-HCl, pH 7.5) for 1 minute, and then to 3MM soaked in Sx SSC: 250 mM Ammonium Acetate for 1 minute. The

3x SSC; 250 mm Ammonium Acetate for 1 minute. The filters were then air dried before baking in an 80°C vacuum oven for 30 minutes.

The oligonucleotide probe was prepared by combining the following:

10 2 µl oligonucleotide (1 pmoles/µl)

2 ul %-32P ATP (3000 Ci/mmole) (Amersham International plc)

 $2~\mu\dot{1}$ 10 x Kinase buffer (0.5 M Tris-HCl, pH 7.5; 100 mM Magnesium Chloride; 10 mM DTT)

15 12 µl Water

2 µl Polynucleotide Kinase (20 Units)

This was incubated at 37°C for 1 hour.

Hybridization was performed in the Techne HB-1 Hybridiser. The baked filters were pre-hybridized at 20 37°C in 40 ml of Hybridization Buffer (10 ml 100 mt Sodium pyrophosphate; 180 ml 5.0 M Sodium chloride; 20 ml 50x Denharts Solution: 90 ml 1.0 M Tris-HCl, pH 7.5; 24 ml 250 mt EDTA; 50 ml 10% NP40; made to 1 litre with water; 60.3 mg rATP; 200 mg yeast RNA (Sigma), for 15

25 minutes before the addition of the 20 µl of the kinased oligo. The filters were incubated at 37°C for at least one hour, and then washed 3 times with 50 ml of 6x SSC at 37°C for 10 minutes (low stringency wash). Filters were air dried, covered with Saran wrap and exposed overnight

30 with Kodak X-AR film.

Selection of fd-CAT2 scFv D1.3 from fd-CAT2 TPB4

Figure 39, summarizes the results from panning experiments using a mixture of the high affinity fd-CAT2 scFv Dl.3 phage (Kd-2 nM) and the fd-CAT2 TPB4 construct

35 (Kd-52 nM).

At a coating concentration of 3000 $\mu g/ml$ Lysozyme, little or no enrichment could be obtained. It was however, possible to get enrichment for the scPv Dl.3 phage when a lower concentration of Lysozyme was used for

40 coating the plates. The best enrichment value obtained was from 1.5% fd-CAT2 scFv Dl.3 in the starting mixture, to 33% fd-CAT2 scFv Dl.3 in the eluted faction, on a plate coated overnight with 30 µg/ml Lysozyme.

Selection of fd-CAT2 serv D1.3 from fd-CAT2 TPB1

Enrichment for the high affinity scFv Dl.3 phage over the fd-CAT2 TPB1 phage (Kd-13) nM, could only be shown from experiments where the plates had been coated overnight with low concentrations of Lysozyme, as shown in Figure 40.

50 In summary, single chain Fv versions of a series of humanized D1.3 antibodies have been constructed in phage fd-CAT2. By affinity selection of fd-CAT2 phage

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mixtures, by panning in small petri dishes, it was shown that the high affinity scFv D1.3 phage, could be preferentially selected for against a background of lower affinity scFv HuD1.3 phage.

Example 36 Expression of Catalytically Active Staphylococcal Nuclease on the Surface of Bacteriophage fd

Examples 11 and 12 showed that alkaline phosphatase from E.coli can be expressed as a catalytically active enzyme on the surface of bacteriophage fd. Here we show 10 that Staphylococcal nuclease can also be expressed in a catalytically active form suggesting that this methodology may be general.

The gene for the enzyme Staphylococcal nuclease (SNase) was amplified from M13 mp18 - SNase (Neuberger, 15 M.S. et al Nature 312 604-608, 1984) by PCR using primers with Internal ApaLI (5'-

GGAATTCGTGCACAGAGTGCAACTTCAACTAAAAAATTAC-3') and NotI

(5"-

GGGATCCGCGGCCGCTTGACCTGAATCAGCGTTGTCTTCG-3') restriction sites, cloned into phage vector fd-CAT2 after digestion with Apall-Notl restriction enzymes and the nucleotide sequence of the SNase gene and junctions with gene III checked by DNA sequencing. The fd-tet-SNase phage was prepared from the supernatant of infected E.coli TG1 cultures by three rounds of PEG precipitation, and the fusion protein demonstrated by SDS-gel electrophoresis and Western blotting using rabbit anti-g3p antiserum (Prof. I. Rasched, Konstanz) and peroxidase-labelled goat anti-rabbit antibodies (Sigma) (Fig. 41) as described in As well as the fusion protein band example 27. (calculated Mr 59749, but runs at a higher position due to the aberrant g3p behaviour), a smaller (proteolytic ?)

product is seen. The fusion protein was shown to be catalytically 35 active by incubation of the fd-tet-SNase phage (4 x 109 tetracyclin resistant colonies [TU]) with single stranded DNA (1 µg) for 1 hr at 37°C in the presence of Ca2+, and analysis of the digest by agarose gel electrophoresis

(Figure 42). Nuclease activity was not detected with the parent fd-CAT2 (2 x 10^{10} TU) phage alone or after three 40 rounds of PEG precipitation of mixtures of fd-CAT2 (2 x 1010 TU) with SNase (0.7 pg). Thus the nuclease activity results from the display of the enzyme on the surface of the phage and not from co-precipitated or soluble SNase 45

set free by degradation of the fusion protein. nuclease activity of fd-tet-SNase (Figure 42) lies in the same order of magnitude, (2 x 108 TV and assuming three copies of SNase per TU) as an equimolar amount of SNase (0.03 ng or 109 particles), and like the authentic SNase 50 was dependent on Ca2+, since incubation with 40 mm MgCl2 and 25 mm EGTA blocked activity (not shown).

Example 37: Display of the Two Aminoterminal Domains of Human CD4 on the Surface of fd Phage

The protein CD4, a member of the immunoglobulin superfamily, is a cell surface receptor involved in MHC class II restricted immune recognition. It is also recognised by the protein gp120 derived from the human immunodeficiency virus (AIDS virus). The first two domains (named V1 and V2, residues 1-178) of the surface antigen CD4 were amplified from pUCl3-T4 (gift from T. 10 Simon) containing the human cDNA of CD4, by PCR using primers with internal ApaLI (5'-GGA ATT CGT GCA CAG AAG AAA GTG GTG CTG GGC AAA AAA GGG G-3') and NotI (5'-GGG ATC CGC GGC CGC AGC TAG CAC CAC GAT GTC TAT TTT GAA CTC-3') restriction sites. After digestion with these two enzymes, the PCR-product was cloned into fdCAT2, and the 15 complete nucleotide sequence of the CD4-VIV2 DNA and junctions with gene III checked by dideoxy sequencing using oligonucleotides fd-seql (5'-GAA TTT TCT GTA TGA GG), CD4-seq1 (5'-GAA GTT TCC TTG GTC CC-3') and CD4-seq2 (5'-ACT ACC AGG GGG GCT CT-3'). In the same way, a fd-20 CD4-V1 version was made, linking residues 1-107 to the Nterminus of gene III, using previously mentioned primers and oligonucleotide 5'-GGG ATC CGC GGC CGC GGT GTC AGA GTT GGC AGT CAA TCC GAA CAC-3' for amplification, FCR conditions and cloning were essentially as described in 25 example 15 except that digestion was with ApaLI and NotI (used according to the manufacturers instructions).

Both fd-CD4-V1 and fd-CD4-V1V2 phages were prepared from the supernatant of infected E.coll TG1 cultures by three rounds of PEG precipitation, thereby concentrating the sample 100-fold for ELISA analysis. The fusion protein was detected in a Western blot (results not shown) with a rabbit anti-gene III antiserum, and revealed bands of the expected size

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Binding of the CD4 moiety to soluble gp120 (recombinant HIV-IIIB gp120 from CHO cells, ADP604, obtained from the Aids Directed Programme, National Institute for Biological Standards and Controls, South Mimms, Potters Bar, UK) was analysed in an ELISA, using 5 µg/ml gp120 for costing (overnight, in PBS). Anti-M13 antiserum was used to detect bound phage; all other conditions were as in Example 9. Figure 43 shows the ELISA signals of wild-type phage (fd-tet) and both CD4phages. Both CD4-phages can bind gp120, but fd-CD4-V1V2 binds much stronger to gp120 than fd-CD4-V1. The binding competitors, soluble CD4 (recombinant soluble CD4 from Baculovirus, ADP 608; from the AIDS Directed Programme) (25 µg/ml) or soluble gp120 (20 µg/ml), added together with the 50 µl phage stock sample during the ELISA. decreased the signal to background level. These results indicate that phage binding to gp120 is mediated by the CD4 molecule displayed at its surface,, and that binding

PCT/GB91/01134

is stronger when the two aminoterminal domains of CD4 are presented.

Thus, CD4 is a cell surface receptor molecule which is active when displayed on bacteriophage fd. Like the PDGF-BB receptor, the functional display of which is described in examples 15 and 16, CD4 is a member of the immunoglobulin superfamily and this result suggests that this class of molecule may be generally suitable for display on the surface of phage.

Example 38 Generation and Selection of Mutants of an 1.0 Anti-4-hydroxy-3-nitrophenylacetic acid (NP) Antibody

expressed on Phage using Mutator strains

It will sometimes be desirable to increase the diversity of a pool of genes cloned in phage, for example a pool of antibody genes, or to produce a large number of 15 variants of a single cloned gene. There are many suitable in vitro mutagenesis methods. However, an attractive method, particularly for making a more diverse population of a library of antibody genes, is to use This has the advantage of generating mutator strains. 20 very large numbers of mutants, essentially limited only

by the number of phage that can be handled. The phage display system allows full advantage to be taken of this

number to isolate improved or altered clones.

Nucleotide sequences encoding an antibody scFv 25 fragment directed against 4-hydroxy-3-nitrophenylacetic acid (NP), scFvB18, derived as in example 14 from a monoclonal antibody against NP were cloned into fdCAT2 using ApaLI and NotI restriction sites as in example 11

to create fdCAT2scFvB18 or into fdDOGKan (fdCAT2 with its 30 tetracycline resistance gene removed and replaced by a kanamycin resistance gene) using PstI and NotI restriction sites to create fdDOGKanscFvB18 or into the phagemid vector pHEN1 using the restriction sites SfiI and NotI as a fusion protein with gene III to create 35

pHEN1scFv818.

The following mutator strains (R. M. Schaaper & R.L. Dunn J. Mol. Biol. 262 1627-16270, 1987; R. M. Schaeper Proc. Natl. Acad. Sci. U.S.A. 85 8126-8130 1988) were

40 used:

NR9232: ara, thi, mutD5-zaf13::Tn10, prolac, F'prolac NR9670: ara, thi, azi, mutTl, leu::TnlO, prolac NR9292: ara, thi, mutH101, prolac, F'prolac

NR9084: ara, thi, mutTl, azi, prolac, F'prolacI"Z"/\M15 45 NR9046: ara, thi, supE, rif, nalA, metB, argE(am), prolac, F'prolac

were kind gifts of Dr. R. M. Schaaper (Department of Health & Human Services, N1H, PO Box 12233, Research

Triangle Park, N.C. 27709) 50 NR9046mutD5: NR9046 mutD5::Tn10 NR9046muttl: NR9046 muttl::Tn10

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were constructed by P1 transduction according to standard procedures. Mutator strains were transfected with fdCAT2scFvB18 of fdDOGKanscFvB18 and transfectants selected for antibiotic resistance. Transfectants were grown for 24h at 37°C before mutant phage was harvested by PEG precipitation. The mutant phage were selected on a lml NIP (4-hydroxy-3-iodo-5nitrophenylacetic acid)-BSA-Sepharose affinity column (prepared according to the manufacturers instructions) prevashed with 200ml of PBS and blocked by 20ml MPBS. Phage were loaded on the column in 10ml MPBS and unbound material reapplied to ensure complete binding. The column was subsequently

washed with 10ml of MPBS and 500ml of PBS. Phage bound to the affinity matrix was eluted with 5 column volumes of 0.33 mM NIP-Cap (example 48).

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Phage eluate was incubated for 30min to 1h with log phase (2x108 cells/ml) E.coli mutator strains without antibiotic selection. The infected cells were then diluted 1:100 in 2xTY and grown for 24h with antibiotic selection (15µg/ml tetracyclin or 30µg/ml kanamycin for fdCAT2scFvB18 or fdDOGKanscFvB18 respectively). Phage from this culture was used for another round of affinity selection and mutation.

Binding of phage antibodies was assayed by ELISA as 25 in example 9 except that ELISA plates were costed with NIP-BSA (4-hydroxy-3-iodo-5-nitrophenylacetyl-BSA; 0.4 Culture supernatants were prepared following growth in Cellwells as described in example 21 and 20µ1 of culture supernatant was added to each well diluted to

30 200ul with MPBS.

Phage samples giving signals in ELISA of more than twice the background were tested ELISA as above for nonspecific binding against lysozyme, BSA or Ox-BSA (example Specificity for NIP was further confirmed by an ELISA in which serial dilutions of NIP-CAP were added 35

together with phage antibodies. Addition of increasing concentrations of NIP-CAP reduced the ELISA signal to the background level.

Phage giving positive signals in ELISA were sequenced and 2 different mutants were subcloned into 40 pHEN1 phagemid and transformed into HB2151 for soluble expression and TG1 for phage display (example 27).

For expression of soluble scrv fragments, transformants in E.coli HB2151 were grown at 37°C in 1 litre 2xTY, 0.2% glucoe, 0.1mg/ml ampicillin to an 0D600 of 1 and expression of soluble scFv fragments induced by adding IPTG to 1mM. Cultures were shaken at 30°C for

Soluble scFvB18 was concentrated from crude 50 bacterial supernatant in a FLOWGEN ultrafiltration unit to a volume of 200ml.

The concentrate was passed two times over a 2ml

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column of NIP-BSA-Sepharose prewashed with 200ml of PBS. The column was washed with 500ml of PBS and 200ml of 0.1M Tris pH7.5, 0.5M NaCl and phage antibodies eluted with 50mM Citrate buffer pH2.3. The eluate was immediately neutralised with 1MTris pH8. The eluate was dialysed against two changes of 1 litre PBS, 0.2mM EDTA. Precipitated protein was removed by centrifugation at 10000g and protein yield was determined by measuring the absorbance at 280nm of the supernatant.

After 4 rounds of mutation and selection, isolated

clones were screened and in one or two rare examples strongly positive ELISA signals were obtained from phage antibodies derived from the mutation of each of fdCAT2scFvB18 and fdDOGKanscFvB18 in the ELISA. The ELISA conditions were such that the parent phage fdCAT2scFvB18 only generated weak signals. These phage antibodies giving strongly positive ELISA signals were enriched in further rounds by a factor of roughly 2.5 per Forty phage antibodies giving strongly positive round. signals were sequenced and they each displayed single mutations in six different positions in the scFvBl8 nucleotide sequences, five of which reside in the light chain. More than 70% of the mutations occurred at positions 724 and 725 changing the first glycine in the J segment of the light chain (framework 4) to serine (in 21 The mutations found

cases) or aspartate (in 3 cases). The sequence of scFvB18 is shown are shown in Table 9. in Figure 44. The nucleotide sequences encoding the scFv fragments

as above.

of a framework mutant with the above glycine to serine mutation, as well as a mutant where Tyr in the CDR3 of the light chain had been mutated to aspartate, were amplified by PCR from the phage antibody clones and subcloned into pHEN1 phagemid (essentially as in example This avoids possible problems with geneIII The same mutations caused by the mutator strains. pattern of ELISA signals was seen when the mutants were displayed on phage following rescue of the phagemid with helper phage (as described in example 25) as when the mutants were assayed when expressed from the phage genome

The scFv fragments from scFvBl8 and the scFv fragments containing the glycine to serine and tyrosine to aspartate mutations respectively were expressed in solution (following transformation into E.coli HB2151 as They showed no differences in in example 27) at 30°C. the ELISA signals between wild-type B18 and the framework mutant. The signal obtained from the phage antibody with the Tvr mutated to aspartate in CDR3 of scFvB18 was about Expression yields were found to be 10x stronger. comparable as judged by Western blotting using an antiserum raised against g3p (as described above).

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Affinity measurements were performed using fluorescence quenching as described in example 23. Affinity measurement of affinity purified scFv fragments however showed scFvBl8, and the scFvBl8 (Gly->Ser) and scFvB18(Tyr->Asp) mutants all to have a comparable affinity of 20nm for NIP-CAP.

A Western blot using an anti-geneIII antibody showed framework mutant had suffered significantly less

proteolytic cleavage than scFvB18. 1.0

Hence, the use of mutator strains generates a diverse range of mutants in phage antibodies when they are used as hosts for clones for gene III fusions. this case some of the clones exhibit higher ELISA signals probably due to increased stability to proteolyic attack. 15 The mutator strains can therefore be used to introduce diversity into a clone or population of clones. diversity should generate clones with desirable characteristics such as a higher affinity or specificity. Such clones may then be selected following display of the

proteins on phage. Example 39 Expression of a Fv Fragment on the Surface of Bacteriophage by Non-Covalent Association of VH and VL domains

This example shows that functional Fv fragments 25 can be expressed on the surface of bacteriophage by noncovalent association of VH and VL domains. One chain is expressed as a gene III fusion and the other as a soluble polypeptide. Thus Fv fragments can be used for all the strategies discussed for Fab fragments including dual 20 combinatorial libraries (example 26).

A useful genetic selection system for stably associated Fv fragments could be established if the expression of Fv fragments as fusion proteins on the phage surface would be possible such that one V domain is fused to the gene III protein and the other V domain is expressed separately in secreted form, allowing it to associate with the V domain on the fusion protein provided the interaction strength is sufficiently high. This idea was tested in a model experiment using the V domains from the anti-hen egg lysozyme antibody D1.3 by fusing the D1.3 VK gene to gene III and separately expressing the D1.3 VH domain.

Experimentally this was achieved as follows: The vector fd-DOG1 was digested with the restriction enzymes PstI and Xhol. From the Fv expression plasmid pSW1-VHD1.3-VKD1.3myc version 3/pUCl19 (Ward et al., 1989 supra) a Pst 1/Xho I-digested restriction fragment was isolated that carries the VH domain coding sequence

(terminated by 2 stop codons), a spacer region between VH 50 and VK genes including a ribosome-binding site for expression of the VK gene, a pelB leader sequence, and, following in frame, the VK gene. This fragment was

cloned into the digested fd-DOG vector to generate the construct fd-tet Fv D1.3. As shown on the map in Fig. 45. the dicistronic VH/VK-gene III operon is transcribed from the gene III promoter: secretion of the VH domain is achieved by the gene III protein leader, secretion of the VK-geneIII fusion protein by the pelB leader sequence. For control purposes a second construct with the name fdtet Fv D1.3 (\(\sigma\si described above: the VH used in this construct carries an insertion of a 200 bp fragment in the Sty I restriction 10 site at the junction of VH CDR 3/FR4, thus interrupting the VH with several in frame stop codons. It is known from previous work that this insertion sufficiently disrupts the VH structure to abolish binding to the antigen lysozyme when expressed either as a soluble Fy or 15 single-chain Fv fragment or as a single-chain Fv fragment on phage surface. This construct was used as a control. TGI bacteria carrying either the fd-tet Fv D1.3, fd-tet Fv D1.3 ((S-Stuffer) or as single-chain wild-type control fd-tet scFv Dl.3 plasmids were grown in liquid culture (medium ZxTy containing 15 µg/ml tetracycline) for 24h to 20 produce phage particles in the supernatant. removal of bacterial cells by centrifugation the phage titer in the supernatants was determined by re-infecting exponentially growing TG1 cells with dilutions of the 28 supernatants and scoring tetracycline-resistants colonies after plating on tetracycline-plates. The infectious phage titers achieved were lx10^{LL} tetR transducing units/ml for the single-chain wild-type control fd-tet SCFV D1.3 and 2x1010 tetR transducing units/ml for Fv phage constructs fd-tet Fv D1.3 and fd-tet Fv D1.3 (\(\Delta \)

Stuffer). ELISA of hen egg lysozyme was performed as in example 2. The results are shown in Fig. 46. Phage derived from

bacteria carrying and expressing the Fv construct fd-tet 35 Fv Dl.3 bind to the immobilised hen egg lysczyme, and when taking the phage titer into account, indeed apparently better than the single-chain FV bearing phages produced by fd-tet scFv D1.3 carrying bacteria. specificity of the reaction and the requirement for a functional VH domain is demonstrated by the fd-tet FV 40 D1.3 (△S-Stuffer) control in which disruption of the VH

domain and consequently of the Fv fragment association eliminates binding to lysozyme.

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As a final control of the expected structure of the VK/geneIII fusion protein a Western Blot was carried out. 20 µl of phage suspensions concentrated 100 fold by two sequential precipitations with PEG were applied to a 10% SDS-PAGE gel. electrophoretically separated and then transferred to a PVDF membrane (Immobilon, Millipore) in 50 a semi-dry Western transfer apparatus (Hoefer). Remaining binding sites on the filter were blocked by lh incubation with 3% BSA in PBS, and detection of the gene III protein accomplished by incubation with a 1:1000 diluted rabbit anti-geneIII antiserum for 2h, neveral washes in PBS/0.1% Tween 20, incubation with peroxidase-conjugated goat anti-rat immunoglobulin antibodies, washes and development with the chromogenic substrate diaminobenzidine/CoCl2/0.03% H2O2. The Fv phage fd-tet Fv D1.3 yields a band for the gene III fusion protein (data not shown), that is intermediate in size between the bands obtained for a wild-type gene III protein from fd-DOG1 and the scFv-gene III fusion protein from fd-tet scFv D1.3, thus proving the presence of a single immunoglobulin domain covalently fused to the gene III

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product int he Fv phage.

In summary, Fv-gene III fusions in which one V domain is fused to the gene III protein and the other V domain associates non-covalently can be presented in functionally active form on the surface of filamentous phage. This opens the possibility to genetically select for stably associated Fv fragments with defined binding specificaties from V gene libraries expressed in phages.

Example 40 A PCR Based Technique for one step Cloning of Human V-genes as Fab Constructs

This example describes a PCR based technique to assemble human Fabs by splicing together the heavy and light chain DNA with a separate piece of 'linker' DNA. A mixture of universal primers is used which should make the technique applicable to all human V-genes.

The general technique for PCR assembly of human Vgenes to create a Fab construct is described. The
efficiency of this technique was assessed by
"assembling", cloning and expressing a human anti rhesusD(Rh-D) Fab from a IgG-K monoclonal hybridoma. We also
demonstrate the potential to rescue human monoclonal
antibodies from polyclonal cell populations by
assembling, cloning, expressing and isolating an IgGlambda monoclonal anti-Rh-D Fab from a polyclonal
lymmoblastic cell line (LCL).

The overall strategy for the PCR assembly is shown in fig.47 and is described in more detail below. For Fab assembly, the VH-CHI and VK-CK or V lambda-C lambda light chains are amplified from first strand cDNA and gel purified. Heavy and light chain DNA are then combined together with linker DNA and flanking oligonuclectides in a new PCR reaction. This results in a full length Fab construct since the 5' end of the linker DNA is complementary to the 3' end of the CHI domain and the 3' end of the linker is complementary to the 5' end of the light chain domain. The linker DNA contains terminal residues of the human CHI domain, the bacterial leader sequence (pelB) for the light chain and the initial residues of the VK or V lambda light chain (fig.2).

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Finally, after gel purification, the Fab construct is reamplified with flanking oligonucleotides containing restriction sites for cloning.

Oligonucleotide primers: In order to develop the PCR cloning of human V genes it was necessary to design a new

range of human specific oligonucleotide primers.

The PCR primers at the 5' end of the VH and VK and Vlambda gene exon (BACK primers) are based on sequence data extracted from the Kabat database, (Kabat, E.A. et al, Sequences of Proteins of Immunological Interest. 4th Edition. US Department of Health and Human Services. 1987) the EMBL database, the literature (Chuchana, P., et al, Eur J. Immunol. 1990. 20:1317) and unpublished data. The sequence of the VH, VK and Vlambda primers are given in table 1. In addition, extended VH primers with Sfil sites at the 5' end were also designed (Table 10) for

adding a restriction site after assembly.

Table 10 also shows the 3' primers (FORWARD primers) designed for the PCR based cloning of human V genes. There are two sets of these depending on whether a Fab or 20 scFv is to be produced. For Fab assembly, the forward primer was based at the 3' end of the CH1 domain, CK domain and Clambda domain. In addition, the CK and C2 FORWARD primers were also synthesized as extended versions with Notl sites at their 5' ends. 25

Primers complementary to the CH1 forward primers and the VkK and V lambda back primers were synthesized to permit generation of linker DNA by PCR amplification of a plasmid template containing the Fab linker (Table 10). To ensure adequate amplification, the primers were

30 extended into the actual linker sequence.

A RNA preparation

This is essentially the same as described in Example 14, but using material of human origin. In the results given in this example human hybridoma and human polycional lymphoblastic cell lines were used.

B cDNA preparation

Approximately 4µg of total RNA in 20ul water was heated at 65°C for 3 minutes, quenched on ice and added to a 30 ul reaction mixture resulting in a 50ul reaction 40 mixture containing 140mM KC1, 50mM Tris, HC1 (pH8.1 @ 42°C), 8mm MgCl2, 10mm DTT, 500uM decxythymidine triphosphate 500 um deoxycytosine triphosphate, 500 um decxyadenosine triphosphate and 500 uM decxyguanosine triphosphate, 80 units of human placental RNAse inhibitor 45 and 10pmol of the appropriate Forward primer (HulgG1-4CHIFOR, HulgMFOR, HuCKFOR, HuCLFOR). Two ul (50 units) of avian myeloblastosis virus (AMV) reverse transcriptase was added, the reaction incubated at 42°C for 1 hour, heated to 100°C for 3 minutes, quenched on ice and 80

centrifuged for 5 minutes.

C Primary PCRs

For the primary PCR amplifications, an equimolar mixture of the appropriate family based BACK and FORWARD primers was used. (See specific examples 40a and 40b given later in this example). A 50ul reaction mixture

was prepared containing 5ul of the supernatant from the cDNA synthesis, 20 pmol total concentration of the FORWARD primers, 250 uM dNTPs, 50mM KC1, 100mM Tris. HC1 (pH 8.3), 1.5 mM MgCl2, 175ug/ml BSA and 1ul (5 units) Thermus aquaticus (Taq) DNA polymerase (Cetus.

10 Emeryville, CA). The reaction mixture was overlaid with paraffin oil and subjected to 30 cycles of amplification using a Techne thermal cycler. The cycle was 94°C for 1 minute (denaturation), 57°C for 1 minute (annealing) and 72°C for 1 minute (extension). The product was analyzed 15 by running 5ul on a 2% agarose gel. The remainder was extracted twice with ether, twice with phenol/chloroform, ethanol precipitated and resuspended in 50ul of H20.

D Preparation of linker

To make the Fab linker DNA, 13 separate PCR 20 reactions were performed using HulgG1-4CH1FOR and each of the reverse VK or V lambda oligonucleotides. template was approximately lng of pJM-1Fab D1.3 (fig.48) The PCR reaction reagents were as described above and the cycle was 94":1 min, 45":1min and 72":1 min. The linkers 25 were analyzed on a 4% agarose gel, purified on a 2%

agarose gel, eluted from the gel on a Spin-X column and ethanol precipitated.

E Assembly PCRs

For PCR assembly of a human Fab approximately lug of 30 a primary heavy chain amplification and lug of a primary light chain amplification were mixed with approximately 250ng of the appropriate linker DNA in a PCR reaction mixture without primers and cycled 7 times (94°: 2 min.

72°:2.5 min) to join the fragments. The reaction mixture was then amplified for 25 cycles (94°:1 mi, 68°-72°:1 35 min, 72°:2.5 min) after the addition of 20 pmol of the appropriate flanking BACK and FORWARD primers.

F Adding Restriction Sites

The assembled products were gel purified and 40 reamplified for 25 cycles (94°:1 min, 55°:1 min, 72°: 25min) with the flanking oligonuceotides containing the appended restriction sites. PCR buffers and NTPs were as described previously.

Specific examples of PCR assembly of human immunoglobulin 45

genes

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a. PCR assembly of a Fab from a human hybridoma: the human monoclonal anti Rh-D cell lines Fog-1 (IgG-k) was derived from EBV transformation of the PBLs of a Rh-D negative blood donor immunized with Rh-D positive blood and has been previously described (Melamed, M.D., et al., J. Immunological Methods. 1987. 104:245) (Hughes-Jones N.C., et al., Biochem. J. 1990. 268:135) (Gorick, B.D. et

al., Vox. Sang. 1988. 55:165) Total RNA was prepared from approximately 10^7 hybridoma cells. First strand cDNA synthesis was performed as described above using the primers HulgG1-4CH1FOR and HuCKFOR. Primary PCRs were performed for the VH-CH1 using a mixture of the 6 HuVHBACK primers and HulgGl-4CGlFOR and for the VK-CK using a mixture of the 6 HuVKBACK primers and HuCKFOR. A Fab construct was assembled as described above, restricted with Sfil and Notl, gel purified and ligated into pJM-1Fab Dl.3 restricted with Sfil and Notl. The 10 ligation mixture was used to transform competent E.coli E.M.G. cells. Ninety-six clones were toothpicked into media in microtitre plate wells, grown to mid-log phase at 30°C and then expression of the Fab was induced by heat shocking at 42°C for 30 min followed by growing for 15 4 hours at 37°C. The ninety-six clones were then screened for anti-Rh-D activity as described below. b. assembly of human Fabs from a polyclonal (LCL); A polyclonal LCL "OG" was derived from EBV transformation of approximately 107 peripheral blood lymphocytes (PBLs) from a Rh-D negative donor immunized with Rh-D positive red blood cells. The cells were plated at a concentration of approximately 105 cells per well. Positive wells were identified by screening the cells harvested and then subcloned once. Typing of the well 25 indicated that an IgG-lambda antibody was being produced. At this stage, total RNA was prepared from approximately 100 cells. First strand cDNA synthesis was performed as described above using the primers HulgGl-4CGlFOR and HuCLFOR. Primary PCRs were performed for the VH-CH1 using a mixture of the 6 HuVHBACK2 primers and HulgG1-4 30 CCIFOR and for the V lambda-C lambda using a mixture of the 7 HuV BACK primers and HuC FOR. Restriction, cloning and screening proceeded as described. To determine the diversity of the clones, the VH and V 35 lambda genes of 15 clones were PCR amplified, restricted with the frequent cutting restriction enzyme BstNl and analyzed on a 4% agarose gel (see example 20). Assay for anti-Rh-D activity and demonstration of specificity: A 5% (vol/vol) suspension of either Rh-D 40 positive (OR2R2) or Rh-D negative (Orr) erythrocytes in phosphate buffered saline (PBS, pH 7.3) were incubated with a papain solution for 10 min at 37°C. The erythrocytes were washed three times in PBS and a 1% 45

with a paper of solution for the times in PBS and a 1% erythrocytes were washed three times in PBS and a 1% (vol/vol) suspension of erythrocytes was made up in PBS supplemented with 1% (vol/vol) of bovine serum albumin (BSA). Fifty ul of a paper treated erythrocyte suspension and 50ul of phage supernatant were placed in the wells of round bottom microtitre plates and the plates were placed on a Titertek plate shaker for 2 min. After 15 min incubation at 37°C 100 ul of PBS/BSA was added to each well. The plates were centrifuged at 200 g

for 1 min and the supernatant was discarded. erythrocytes were resuspended in the remaining FBS/BSA and the Fab fragments were crosslinked by addition of the 9E10 monoclonal antibody (50ul a lug/ml solution in PBS/BSA) directed against the myc peptide tag (Ward, E.S., et al., Nature 1985. supra). The plates were placed at room temperature (RT) until sedimentation had occurred. Agglutination of erthrocytes caused a diffuse button of erythrocytes and the results were evaluated macroscopically. Specificity was confirmed with a 10 standard prepapainized (as above) panel of 9 erythrocyte suspensions in PBS (all suspensions blood group O, 4 D positive and 5 D negative) known to have homozygous expression of all the clinically relevant erythrocyte blood group alloantigens. The number of copies of the D 15 antigen on the D positive cells varied between 10,000 and 20,000 per erythrocyte depending on the Rh genotype. Briefly, 50 ul phage supernatant in PBS supplemented with 2% (vol/vol) skimmed milk was mixed with 50 ul of a 2% erythrocyte suspension in PBS in glass tubes and 20 incubated for 15 min at 37°C. After one wash with PBS/BSA the erythrocytes were pelleted and resuspended in 50 ul donkey anti-human lambda light chain (Sigma L9527,

diluted 1:40 in PBS/BSA). The tubes were centrifuged for 25 1 min at 200g and agglutination was read macroscopically using "tip and roll" method.

Results

30 a PCR assembly of a Fab from a human hybridoma: A single band of the correct size was obtained after amplification. Thirty-eight of 96 clones (40%) screened specifically agglutinated Rh-D positive but not Rh-D negative red blood cells. The results demonstrate a high

35 frequency of successful splicing in the assembly process and the potential of this technique for one step cloning of human hybridomas.

b Assembly of human Fabs from a polyclonal lymphoblastic

cell line (LCL): Analysis of the diversity of the clones of indicated that 3 different heavy chain families and 2 different light chains families were present. Five anti-Rh-D specific clones were identified out of 96 screened. The VH and VA chains had identical nucleotide sequences

in each clone and were typical of anti-Rh-D V-genes (unpublished results). The results demonstrate the potential of this technique to assemble, clone and isolate human antibody fragments from polycional cell populations (see also section on isolation of specific

binding activities from an 'unimmunized' human library (examples 42 and 43).

Example 41
Selection of Phage Displaying a Human Fab Fragment

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directed against the Rhesus-D Antigen by binding to Cells

displaying the Rhesus D Antigen on their Surface A large number of important antigens are integral components of cell surface membranes, i.e. they are cell surface antigens. These include tumor specific antigens and red and white blood cell surface antigens. instances, it would be important to isolate antibodies against these antigens. For example, antibodies directed against the rhesus-D (Rh-D) antigen on red blood cells are used both diagnostically and therapeutically. Many 10 of these antigens are difficult to purify and some, like Rh-D, are not biologically active when isolated from the Thus, it would be useful to be able to membrane. affinity purify antibody fragments displayed on the surface of bacteriophage directly on cell surface 15 antigens. To test the feasibility of affinity purification on cell surface antigens, the anti-Rh-D

purification on cell surface antiques, case antihuman monoclonal antibody Fog-8 was displayed as a Fab fragment on the surface of bacteriophage fd. The displayed Fog-8 Fab fragment bound antigen as determined by agglutination assay and could be affinity purified on the basis of its binding on the surface of Rh-D positive

red blood cells but not Rh-D negative red blood cells. Materials and Methods

Construction of a clone encoding an anti-Rh-D Fab fragment in phagemid pHENI and display of the Fab fragment on the surface of bacteriophage fd.

The human hybridoma Fog-B has been previously described (N.C. Hughes-Jones et al Biochem, J. 258 135 (1990). It produces an IgG-1/lambda antibody which binds the Rh-D antigen. RNA was prepared from 107 hybridoma cells using a modified method of Cathala (as described in example 14) and 1st strand cDNA synthesized using specific immunoglobulin heavy and light chain primers (HuVHIFOR [example 40] and HuCA FOR (5'-GGA ATT CTT ATG

ANG ATT CTG TAG GGG CCA C-3') as described in example
14. The VH gene was subsequently amplified from an
aliquot of the lst strand cDNA using HuVHABACK and
HuVHIFOR. The VA gene was amplified using a Waprimer

40 specific for Fog-B (VAFog-B, 5'-AAC CAG CCA TGG CC AGT CTG TGT TGA CGC AGC C-3'). The PCR conditions were as described in example 40. The PCR products were analyzed by running 5ul on a 2% agarose gel. The remainder was extracted twice with ether, twice with phenol/chloroform.

45 ethenol precipitated and resuspended in 50µl of H₂O. The amplified VH DNA was digested with Pstl and BstEII, and the amplified Vh-C. DNA with Nool and EcoRl. The fragments were purified on a 2% agarose gel. extracted using Geneclean, and sequentially ligated into the 50 soluble expression vector pJM-1 Fab Dl.3 (Fig 48). Clones containing the correct insert were initially

Clones containing the correct insert were initially identified by restriction analysis and verified by assay

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of expressed soluble Fab (see example 23 for induction conditions). The Fog-B Fab cassette was amplified from puM-1 by PCR using HuVHABACK-Sfi and Hu CA-Not, digested with the appropriate restriction enzymes and ligated into pHEN1. Clones containing the correct insert were identified initially by restriction analysis and subsequently by assay (see example 25 for induction conditions).

Assay for soluble Fog-8 Fab fragment and phage displayed Fog-8 Fab fragment for anti-Rh-D activity and

documentation of specificity.

Assay of the soluble expressed Fab was performed on unconcentrated E.coli supernatant. Assay of Fog-E displayed on the phage surface was performed on phage that had been concentrated 10 fold by PEG precipitation and then resuspended in PBS. the assays for activity and specificity are as described in example.

Call surface affinity purification of phage

displaying Fog-B anti-Rh-D Fab fragment

20 Purified Fog-B phage was mixed with purified phage Fd-Tet CAT-1 displaying the anti-lysozyme scFv D1.3 (pAbD1.3) in a ratio of approximately 1 Fog-B:50 scFvD1.3. Prepapainized erythrocytes (OR2R2 [Rhesus positive] or Orr [Rhesus negative]) were suspended in PBS 25 supplemented with 2% skimmed milk powder in a concentration of $4\times107/\text{ml}$. One ml of this suspension was mixed with 10^{11} phage suspended in 2 ml of PBS supplemented with 2% skimmed milk and incubated for 30 min at room temperature under continuous rotation. 30 erythrocytes were washed three times with an excess of ice-cold PBS (10 ml per wash) and subsequently pelleted. The phage were eluted from the cells by resuspending in 200 ul of 76 mM citric acid pH 2.8 in PBS for 1 min. The cells were then pelleted by contrifugation for 1 min at 3000 rpm and the supernatant containing the eluted phage 35 was neutralized by adding 200 µl of 240 mM Tris-base, 22mM Disodium hydrogen phosphate in 1% w/vol albumin. Serial dilutions of the eluate was used to infect TG1 cells. Fog-B Fab phage were selected on ampicillin 40 plates and scFvDl.3 phage on tetracycline plates and the titre of each determined prior to selection, after selection on rhesus-D negative cells and after selection on rnesus-D positive cells.

Results
Fog-B Fab fragment displayed on the surface of the phage derived from the phagemid pHEN clone specifically agglutinated rhesus-D positive but not rhesus D-negative red blood cells. Affinity purification of the Fog-I Fab phagemid on Rh-D positive red blood cells resulted in an enrichment from 0:50 to 1500:1 (Fog-B Fab:scFvbl.3), whereas purification on Rh-D negative red blood cells demonstrated essentially no enrichment (10 fold).

RATIO

45

library

TITRE

	Fog-B Fab scFvD1.3 Fog-B FAb/scFvD1.3
5	Prior to selection 1.0 \times 10 ⁸ 5.0 \times 10 ⁹ 1:50 Selection on Rh-D 2.0 \times 10 ⁴ 1.0 \times 10 ⁵ 1:5
	negative cells Selection on Rh-D 6.0 x 10 ⁶ 4.0 x 10 ³ 1500:1 positive cells
10	Example 42 A PCR Based Technique for One Step Cloning of
	Human scFV Constructs Assembly of human scFv is similar to the assembly of mouse scFvs described in example 14. To develop the PCR
15	cioning of human V genes it was necessary to design a new range of human specific oligonucleotide primers (table 10). The use of these primers for the generation of human Fabs is described in example 40. The assembly of human scrys is essentially the same but requires a set of
20	FORWARD primers complementary to the J segments of the VH, VK and V lambda genes. (For Fabs FORWARD primers complementary to the constant region are used.) The J segment specific primers were designed based on the published JH JK and J lambda sequences (Kabat, E.A. et
25	al, Sequences of Proteins of Immunological Interest. 4th Edition. US Department of Health and Human Services. 1987). In addition, a different linker is needed for scrysthm for Table so for human scrys a new set of primers was
30	needed to prepare the linker. Primers complementary to the JH forward primers and the VK and V lambda back primers were synthesized to permit generation of linker DNA by PCR amplification of a plasmid template containing the soft linker (Table 10, Fig. 49). To ensure adequate
35	amplification, the primers were extended into the actual linker sequence. Using these primers to make the scPv linker DNA, 52 separate FCR reactions were performed using each of the 4 reverse JH primers in combination with each of the 13 reverse VK and V lambda
40	oligonucleotides. The template was approximately lng of pSW2scDl.3 (Ward, E.S. 1989 supra) containing the short peptide (Gly4Ser)3 (Huston, J.S. et al., Gene 1989, 77:61) A specific example of PCR assembly of a human scFv
	A CARACA CARACA A CAR

This example describes the generation of a human library of scrys made from an unimmunized human: 500ml of blood, containing approximately 10⁸ B-cells, was obtained from a healthy volunteer blood donor. The white cells were separated on Ficoll and RNA was prepared as described in example 14.

Twenty percent of the RNA, containing the genetic

material from approximately 2 x 107 B-cells, was used for cDNA preparation as described in example 40. Heavy chains originating from IqC and IqM antibodies were kept separate by priming cDNA synthesis with either an IGG 5 specific primer (HulgGl-4CH1FOR) or an IgM specific primer (HulgMFOR). Aliquots of the cDNA was used to generate four separate scFv libraries (IgG-K, IgG-lambda, IgM-K and IgM-lambda) as described in example 40. The resulting libraries were purified on 1.5% agarose,

electroeluted and ethanol pracipitated. For subsequent cloning, the K and lambda libraries were combined giving 10 separate IqG and IgM libraries.

Cloning of the library: The purified scFv fragments (1-

4ug) were digested with the restriction enzymes NotI and 15 either Sfil or Ncol. After digestion, the fragments were extracted with phenol/chloroform, ethanol precipitated. The digested fragments were ligated into either Sfil-NotI or Ncol-Notl digested, agarose gel electrophoresis purified pHEN1 DNA (6ug) (see example 24), in a 100 ul ligation mix with 2,000 U T4 DNA ligase (New England Biolabs) overnight at room temperature. The ligation mix was purified by phenol extraction and ethanol precipitated. The ligated DNA was resuspended in 10 μl of water, and 2.5 µl samples were electroporated into E.coli TG1 (50 μ l). Cells were grown in 1 ml SOC for 1 hr and then plated on 2 x TY medium with 100 μ g/ml ampicillin and 1% glucose (AMP-GLU), in 243 x 243 mm 25

dishes (Nunc). After overnight growth colonies were scraped off the places into 10 ml 2 x TY containing AMP-30 GLU and 15% glycerol for storage at -70°C as a library

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Cloning into Sfil-Not1 and Ncol-Not1 digested pHEN1 yielded libraries of 107 and 2 x 107 clones respectively for the IgM libraries and approximately 5 x 107 clones for each of the two IoG libraries.

Example 43 Isolation of binding activities from a library of scrvs from an unimmunized human

The ability to select binding activities from human 40 antibody libraries displayed on the surface of phace should prove even more important than isolation of binding activities from murine libraries. because the standard way of generating antibodies via hybridoma technology has not had the success with human antibodies that has been achieved with mouse. While in 45 some instances it will be possible to make libraries from immunized humans, in many cases, it will not prove possible to immunize due to toxicity or lack of availability of an appropriate immunogen or ethical considerations. Alternatively, binding activities could be isolated from libraries made from individuals with

diseases in which therapeutic antibodies are generated by

the immune response. However, in many cases, the antibody producing cells will be located in the splean and not available in the circulating pool of peripheral blood lymphocytes (the most easily accessible material for generating the library). In addition, in diseases associated with immunosuppression, therapeutic antibodies may not be produced.

An alternative approach would be to isolate binding activities from a library made from an unimmunized individual. This approach is based on estimates that a primary repertoire of 10' different antibodies is likely to recognize over 99% of epitopes with an affinity constant of 10° M¹ or better. (Pewrelson, A.S. Immunol. Rev. (1989) 110:5). While this may not produce high affinity antibodies, affinity could be boosted by

Rev. (1989) 110:5). While this may not produce high affinity antibodies, affinity could be boosted by mutation of the V-genes and/or by using the isolated VH domain in a hierarchical approach with a library of light chains (or vice versa). In this section, we demonstrate the feesibility of this approach by isolating specific antigen binding activities against three different

20 antigen binding activities against three different antigens from a library of scFvs from an unimmunized human.

Materials and Methods

The generation of the human scFv library used for the isolation of binding activities described in this example is detailed in example 42.

Estimation of diversity of original and selected libraries: Recombinant clones were screened before and after selection by PCR (example 20) with primers LMB3 30 (which sits 5' of the pelB leader sequence and is identical to the reverse sequencing primer (-40 n) of pUC19) and fd-SEQ1 (see example 37) followed by digestion with the frequent-cutting enzyme BstN1. Analysis of 48 clones from each unselected library indicated that 90% of

35 the clones had inset, and the libraries appeared to be extremely diverse as judged by the BstNI restriction pattern.

Rescue of Phagemid libraries for enrichment experiments: To rescue phagemid particles from the library, 100 ml 2 x 40 TY containing AMP-GLU (see example 42) was inoculated with 10⁹ bacteria taken from the library (prepared in example 42) (approx. 10 µl) and grown for 1.5 hr, shaking at 37°C. Cells were spun down (IEC- centrifuge, 4 K, 15 min) and resuspended in 100 ml prewarmed (37°C) 2 x TY-

min) and resuspended in 100 ml prewarmed (37°C) 2 x TY45 AMP (see example 41) medium, 2 x 10¹⁰ pfu of VCS-M13
(Stratagene) particles added and incubated 30 min at 37°
without shaking. Cells were then transferred to 900 ml 2
x TY containing ampicillin (100 µg/ml) and kanamycin (25
µg/ml) (AMP-KAN), and grown overnight, while shaking at

50 37°C. Phage particles were purified and concentrated by three PEG-precipitations (see materials and methods) and resuspended in PBS to 10¹³ TU/ml (ampicillin resistant

clones).

Enrichment for phOx: BSA binders by selection on tubes: For enrichment, a 75 x 12 mm Nunc-immunotude (Maxisorp; Cat. No. 4-44202) was coated with 4 ml ph0x:BSA (1 mg/ml; 33 14 phOx per BSA in 50 mM NaHCO3 pH 9.6 buffer) overnight at room temperature. After washing three times with PBS. the tube was incubated for 2 hr at 37°C with PBS containing 2% Marvel (2% MPBS) for blocking. Following three PBS washes, phagemid particles (1013 TU) in 4 ml of 2% MPBS were added, incubated 30 min at room temperature 10 on a rotating turntable and left for a further 1.5 hours. Tubes were then washed with 20 washes of PBS, 0.1% Tween 20 and 20 washes PBS (each washing step was performed by pouring buffer in and out immediately). Bound phage 15 particles were eluted from the tube by adding 1 ml 100 mm triethylamine pH 11.5 and rotating for 15 min. eluted material was immediately neutralised by adding 0.5 ml 1.0 M Tris-HCl, pH 7.4 and vortexed. Phage was stored at 4°C. Eluted phage (in 1.5 ml) was used to infect 8 ml

20 Éluted phage (in 1.5 ml) was used to infect 8 ml logarithmic growing E.coli TG1 cells in 15-ml 2 x TY medium, and plated on AMP-GLU plates as above yielding on

average 107 phage infected colonies.
For selection of phOx:BSA binders, the rescue-tube

25 enrichment -plating cycle was repeated 4 times, after which phagemid clones were analysed for binding by ELISA. Enrichment for lysozyme binders by panning and on columns: A petri dish (35 x 10 mm Falcon 3001 Tissue culture dish) was used for enrichment by panning. During 30 all steps, the plates were rocked on an A600 rocking plate (Raven Scientific). Plates were coated overnight with 1 ml turkey egg white lysozyme (3 mg/ml) in 50 mM sodium hydrogen carbonate (pH 9.6), washed three times with 2 ml PBS, and blocked with 2 ml 2% MPBS at room 35 temperature for 2 hours. After three PBS washes approximately 10^{12} TU phage particles in 1 ml 2% MPBS were added per plate, and left rocking for 2 hr at room temperature. Plates were washed for 5 min with 2 ml of the following solutions: 5 times PBS, PBS-Tween {0.02% 40 Tween-20), 50 mM Tris-HCl (pH 7.5) + 500 mM NaCl, 50 mM Tris-HC1 (pH 8.5) + 500 mM NaC1, 500 mM Tris-HC1 (pH 9.5) + 500 mM NaCl and finally 50 mM sodium hydrogen carbonate pH 9.6 Bound phage particles were then eluted by adding 1 ml 100 mM triethylamine pH 11.5 and rocking for 5 min 45 before neutralising with 1 M Tris-HC1 (pH 7.4) (as above). Alternatively, 1 ml turkey egg white lysozyme-Sepharose columns were used for affinity purification (McCafferty, J., et al., Nature 1990. 348: 552) Columns were washed extensively with PBS, blocked with 15 ml 2% 60 MPBS, and phage $(10^{12}\ \text{TU})$ in 1 ml 2% MPBS loaded. After washing with 50 ml PBS, 10 ml PBS- Tween (PBS + 0.02%

Tween-20), 3 ml of 50 mM Tris-HCl (pH 7.5) - 500 mM NaCl.

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5 mM Tris-HCl 9pH 8.5) + 500 mM NaCl, 5ml of 50 mM Tris-HCl (pH 9.5) + 500 mM NaCl and finally 5 ml of 50 mM sodium hydrogen carbonate pH 9.6. Bound phage was eluted using 1.5 ml 100 mM triethylamine and neutralised with 1

5 M Tris-HC1 (pH 7.4).

For selection of turkey egg white lysozyme binders, the rescue-tube enrichment-plating cycle or rescue-column-plating cycle was repeated 4 times, after which phacemid clones were analysed for binding by ELISA.

- Rescue of individual phagemid clones for ELISA: Clones resulting from reinfected and plated phage particles eluted after 4 rounds of enrichment, were inoculated into 150 ul of 2 x TY-AMP-GLU in 96-well plates (cell wells, Nunclon), grown with shaking (250rpm) overnight at 37°C.
- Nuncion), grown with snaking (2007pm) overhight at 37 c.

 A 96-well plate replicator ('plunger') was used to inoculate approximately 4 µl of the overnight cultures on the master plate into 200 µl fresh 2 x TY-AMP-GLU. After l hr, 50 µl 2 x TY-AMP-GLU containing 10⁸ pfu of VCS-M13 was added to each well, and the plate incubated at 37°C for 4 for followed by shaking the plate at 37°C for 1 hr. Glucose was then removed by spinning down the cells (4K, 15 min), and aspirating the supernatant with a drawn
- (4K, 15 min), and aspirating the supernatant with a drawn out glass pasteur pipet. Cells ware resuspended in 200 µl 2 x TY-AMP-KAN (Kanamycin 50 ug/ml) and grown 20 hr, 25 sheking 37°C. Unconcentrated supernatant containing
- S shaking 37°C. Unconcentrated supernatant containing phage was taken for analysis by ELISA.
 ELISA
- Analysis for binding to phOx:BSA, BSA or lysozyme was performed by ELISA (see example 9), with 100 µg/ml phOx:BSA or BSA, or 3 mg/ml turkey egg white lysozyme used for coating. Determination of cross reactivity to unrelated antigens with the isolated clones was also determined by ELISA on plates coated with 100 ug/ml of an irrelevant antigen (keyhole limpet haemocyanin (KLH), ovalbumin, chymotrypsinogen, cytochrome C, thyroglobulin, GAP-DH (glyceraldehyde-3-phosphate dehydrogense), or
- trypsin inhibitor).

 Characterization of ELISA positive clones: All antigen specific clones isolated were checked for cross reactivity against a panel of irrelevant antigens as described above. The diversity of the clones was determined by PCR screening as described above and at least two clones from each restriction pattern were

sequenced by the dideoxy chain termination method.

Results

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Isolation and characterization of ph0x:BSA binders:
After 4 rounds of selection, ELISA-positive clones were
isolated for ph0x:BSA. All clones originated from the
for light library. Of 96 clones analysed, 43 clones were
binding to both ph0x:BSA and BSA, with ODs ranging from
0.4 to 1.3 (background 0.125). These clones are

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designated as BSA binders. The binding to BSA seemed to be specific, since none of the 11 clones analysed gave a signal above background when used in an ELISA with KLH, ovalbumin. cnymotrypsinogen. cytochrome C. lysozyme, thyroglobulin. GAP-DH. or trypsin inhibitor. all BSA binding clones had the same BstNI restriction partern, and 14 clones were completely sequenced. Thirteen of the fourteen clones had the same sequence, the VH was derived from a human VH3 family gene and the VL from a human V lambda 3 family gene (Table 1). The other BSA binder was derived from a human VH4 family gene and a human Vk1 derived from a human VH4 family gene and a human Vk1

family gene (data not shown).

One clone was isolated which bound to ph0x:ESA only (OD 0.3), and bound phage could be completed off completely by adding 0.02 mM 4-6-amino-caproic acid methylene 2-phenyi-oxazol-5-one (ph0x-CAP) as a competitor. Also no binding above background could be detected to the panel of irrelevant proteins described above. The sequence revealed a VH derived from a human V lambda 1 VH family gene and a VL derived from a human V lambda 1

family gene (Table 11).

Isolation and characterisation of lysozyme binders: After 4 rounds of selection, 50 ELISA-positive clones were isolated for turkey lysozyme. The majority of the clones, greater than 95%, were from the IgM library. The binding to lysozyme seemed to be specific, since none of the clones analysed gave a signal above background when used in an ELISA with KLH, ovablumin, chymotrypsinogen, cytochrome C, thyroglobulin, GAF-DH, or trypsin inhibitor. The lysozyme binding clones gave 3 different SETNI restriction patterns, and at least 2 clones from

each restriction pattern were completely sequenced. The sequences indicated the presence of 4 unique human VH-VL combinations. (Table 11).

35 Conclusion

The results indicate that antigen binding activities can be isolated from repertoires of scFvs prepared from 1gM cDNA from human volunteers that have not been specifically immunized.

40 Example 44

Rescue of human IgM library using helper phage lacking gene 3 (5g3)

This example describes the rescue of gene 3 fusions from a human library using a helper phage with a gene 3

45 deletion.

100 µl of bacterial stock of the IgM phagemid library prepared as described (example 42), containing 5x10° bacteria, was used to inoculate 100mls of 2xTY medium containing 100µg/ml ampicillin. 2% glucose (TY/Amp/Glu). This was grown at 37°C for 1.5 hours. 10 mls of this culture was added to 90 mls of prewarmed TY/Amp/Glu and infection carried out by adding 10mls of a

(M13K07gIII A No.3) (example 34) and incubating for 1 hour at 37°C without shaking. Preparation of M13K07gIII No.3 was as described in example 34. After centrifugation at 4,000 r.p.m. for 10 minutes the bacteria were resuspended in 100 mls of 2 x TY medium containing 100 µg/ml ampicillin (with no glucose). Titration of the culture at this point revealed that there were 1.9x10d infected bacteria as judged by their ability to grow on plates containing both ampicillin (100µg/ml) and kanamycin 1.0 (50µg/ml). Incubation was continued for 1 hour with shaking before transferring to 2.5 litres of 2xTY medium containing 100µg/ml ampicillin, 50µg/ml kanamycin. contained in five 2.5 litre flasks. This culture was incubated for 15 hours and the supernatant prepared by 15 (10-15 minutes at 10,000 r.p.m. in a centrifugation. Sorvall RC5B centrifuge at 4°C). Phage particles were harvested by adding 1/5th volume of 20% polyethylene glycol, 2.5 M-NaCl, standing at 4°C for 30 minutes and centrifuging as above. The resulting pellet was 20 resuspended in 40mls of 10mM Tris, 0.1mM EDTA pH 7.4 and bacterial debris removed by centrifugation as above. The packaged phagemid preparation was then re-precipitated, collected as above and resuspended in 10mls of 10mM Tris, O.lmM EDTA pH 7.4. The litre of this preparation was 25 4.1x10¹³ transducing units/ml (ampicillin resistance).

Tubes coated with OX-BSA were prepared as described in example 45 for panning the phagemid library from example 42. The rescued library was also panned against tubes coated with bovine thyroglobulin (Sigma). These were coated at a concentration of lmg/ml thyroglobulin in 50mM NaHCO3 pH9.6 at 37°C, overnight. Tubes were blocked with PBS containing 2% milk powder (PBS/M) and incubated with lml of the rescued phagemid library (the equivalent of 250mls of culture supernatant) mixed with 3mls of PBS/M for 3 hours. Washing, elution, neutralisation and infection were as described in example 45.

Results: Panning against oxazalone - BSA

The first round of panning against OX-BSA yielded 2.8x10⁰ phage. A large bacterial plate with 1.4x10⁰ colonies derived from this eluate was scraped into 10mls of 2xxTY, 20% glycerol, shaken for 10 minutes, aliquoted and stored. This was also used to inoculate a frash culture for rescue with M13K07gIII No.3. (Bacteria and rescued phage derived from first round panning against OX-BSA are named OXPANI. Bacteria or rescued phage derived from second and third round pannings are named OXPAN2 and OXPAN3 respectively) Rescue of phagemid with M13K07gIII No.3 after each round of panning was essentially as described above but using 5ml volumes for the initial cultures in TY/Amp/Glu. using 1ml of helper ohage and transferring to 100-500mls of 2xTY medium

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containing 100µg/ml ampicillin, 50µg/ml kanamycin. Second and third round panning steps were as described above for the first round, but using 0.8-1.0mls of 100 fold concentrated phage (the equivalent of 80-100 mls of culture supernatant). The eluate from the second round panning contained 8x10⁸ infectious particles and the eluate from the third round panning contained 3.3x109 infectious particles.

Panning against thyroglobulin

The first round panning against thyroglobulin yielded 2.52x105 infectious particles. Half of the eluate was used to generate 1.26x105 bacterial colonies on a large plate. These colonies were scraped into 10mls of 2xTY, 20% glycerol, shaken for 10 minutes, aliquoted and stored. These bacteria and rescued phage derived from them are termed THYPANI, and used to inoculate a fresh culture for rescue with M13K07qIII No.3 to give a polyclonal rescued phage preparation. Material similarly derived from second and third round pannings are termed THYPAN2 and THYPAN3 respectively. Second and their round pannings with thyroglobulin were as described for second and third round OX-BSA panning. The eluate from the

second round panning contained 8×10^7 transducing units and the eluate from the third round panning contained 6x107 infectious particles.

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ELISA screening of clones derived by panning 40 colonies derived form the third round of panning

against thyroglobulin (THYPAN3) were picked into a 96 well plate and grown overnight at 37°C in 200ul of 30 TY/Amp/Glu. Similarly 48 colonies from two rounds and 48 colonies from three rounds of panning against OX-BSA were grown (OX-PAN2 and OX-PAN3). Polyclonal phage were prepared at the same time. Next day 5µl from each culture was transferred to 100µl of fresh prewarmed TY/Amp/Glu grown for 1.5 hours and M13K07gIII No.3 added 38 (2 x 10^5 infectious phage per well in 100ul of TY/Amp/Glu). these were incubated for 1 hour at 37°C without shaking, centrifuged at 4,000 r.p.m. for 10 minutes, resuspended in 150ul of 2xTY medium containing 40 100µg/ml ampicillin and incubated for a further hour with shaking before adding to 2mls of medium containing 100ug/ml ampicillin, 50µg/ml kanamycin. After overnight

growth the cultures were centrifuged at 4,000 r.p.m. for 10 minutes and the supernatants collected. ELISA plates 45 used to screen THYPAN3 clones were coated at 37°C overnight with 200µg/ml thyroglobulin in 50mM NaHCO3pH9.6. Plates used for OXPAN2 and OXPAN3 were coated at 100µg/ml OX-BSA in PBS at 37°C overnight.

120µl of culture supernatant was mixed with 30µl of 5x PBS. 10% milk powder and incubated at room temperature for 2 hours at room temperature. ELISAs were carried out as described in example 18.

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For thyroglobulin, 18 out of 40 clones were positive (0.3-2.0 O.D. after 30 minutes). (A phage control (vector pCAT3) gave a reading of 0.07 0.D.). addition, positives were also seen on the polyclonal phage preparations THYPAN1 (0.314 0.D.) and THYPAN2 (0.189 0.D.) compared with phage derived from the original non-panned phagemid library (0.069 O.D.). All polyclonal phage were PEG precipitated and used at a 10 fold concentration.

PCR reactions and BstN1 digests were carried out on 10 the positive clones as described above and six different patterns of DNA fragments were obtained showing that at

least six different clones had been isolated.

For OX-BSA after two rounds of panning, 30 of 48 clones were positive by ELISA and after three rounds , 42 15 of 48 were positive. In a separate experiment, positive signal was obtained from the polyclonal phage preparations OXPAN1 (0.988 OD) and OXPAN2 (1.717 OD) compared with phage derived from the original non-panned phagemid library (0.186 O.D.) after 30 minutes.

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Selected clones for thyroglobulin or OX-BSA Selected clones (11 anti-thyroglobulin, 5 anti-OX-BSA) representing each of the different BstNI restriction digest patterns were assayed for binding to a panel of irrelevant antigens. ELISA plates were coated with 25 antigen (100 µl/ml in 50 mM NaHCO3, pH 9.6) by overnight incubation at 37°C. The panel of antigens consisted of keyhole limpet haemocyanin, hen egg lysozyme, bovine serum albumin,ovalbumin, cytochrome c, chymotrysinogen, trypsin inhibitor, GAP-D11 (glyceraldehyde-3-phosphate 30 dehydrogenase), bovine thyroglobulin and oxazolone-BSA. Duplicate samples of phage supernatant (80 µl + 20 µl 5 x PBS, 10% milk powder) were added to each antigen and incubated for 1 hour at room temperature. the ELISA was

carried out as described in example 18. Each of the thyroglobulin specific clones (11 from 11) were positive for thyroglobulin (OD 0.12 - 0.76) but after 60 minutes showed no binding (OD<0.03) to any of the 9 irrelevant antigens. Similarly of the 5 OX-BSA specific clones 3 had an OD 0.07 - 0.52 compared to ODs < 0.02 for the irrelevant antigens. None of the 5 clones

had any binding to BSA alone.

Thus positive clones can be isolated after only two rounds of panning by rescuing with M13K07gIII No.3. In addition there is a greater likelihood with this helper 45 of generating phage particles with more than one intact antibody molecule. This will potentially increase the avidity of phage-antibodies and may enable isolation of clones of weaker affinity.

Example 45: Alteration of fine specificity of scry D1.3 50 displayed on phage by mutagenesis and selection on

immobiliséd turkev lysozyme

The D1.3 antibody binds hen egg lysozyme (HEL) with an affinity constant of 4.5 \times 10 7 M $^{-1}$ whereas it binds turkey egg lysozyme (TEL) with an affinity of $(1\times10^{5}$ M $^{-1}$. (Harper et al (1987) Molecular Immunology 24 99 7 -108, Amit et al (1986) Science 233 2 74 7 -753).

It has been suggested that this is because the glutamine residue present at position 121 of HEL (gln121) is representated by histidine residue at the same position in TEL. Thus mutagenising the Dl.3 antibody residues which interact with gln121 of HEL may facilitate binding to TEL.

According to Amit at al. supra, tyrosine at amino acid position 32, phenylalanine at position 91 and tryptophan at position 92 of the light chain interact with ginl21 of HEL. In addition tyrosine at position 101 of the heavy chain also interacts. None of these residues are predicted to be involved in determining the main chain conformation of the antibody variable regions

(Chothia and Lesk (1987) Journal of Molecular Biology 196, p901-917).

Mutagenesis of pCAT3SCFvD1.3

The cligonucleotides mutt91,92, was prepared too randomise phenylalanine at position 91 (L91) and tryptophan at position 92 (L92) of the light chain. The cligonucleotides mutt32, was prepared to randomise tyrosine at light chain position 32 (L32) and the cligonucleotides mutH101 was prepared to randomise tyrosine at position 101 of the heavy chain (H101). mutt91,92:

30 5' CGT CCG AGG AGT ACT NNN NNN ATG TTG ACA GTA ATA 3' murl32:
5' CTG ATA CCA TGC TAA NNN ATT GTG ATT ATT CCC 3' murl101:

E' CCA GTA GTC AAG CCT NNN ATC TCT CTC TCT GGC 3'

35 N represents a random insertion of equal amounts of A,C,G or T) in vitro mutagenesis of the phagemid vector, pCAT3scFvDl.3 (example 17) with the oligonucleotide mutL91,92 was carried out using an in vitro mutagenesis kit (Amersham). The resultant DNA was transformed by electroporation into TGI cells using a Bio-Rad electroporator. 78,000 clones were obtained and these

were scraped into 15mls of 2xTY/20% glycerol. This pool was called D1.3191192. Single stranded DNA was prepared by rescue with M13KO7 as described in Sambrook et al. 45 1989 supra, and sequenced with the primer FDTSEQ1.

5 1989 supra, and sequenced with the primer FDTSEQ1, using a Sequenase sequencing kit (United States Biochemical Corporation).

This revealed that the DNA had been successfully mutagenised as judged by the presence of bands in all four DNA sequencing tracks at the nucleotide positions encoding 191 and 192. This mutagenised single stranded DNA was subjected to a further round of mutagenesis as

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above using either mutL32 or mutH101 cligonucleotides. Mutagenesis with mutL32 gave rise to 71,000 clones (pool called D1.3L32) while muth101 gave 102,000 clones (pool called D1.3H101). These clones were scraped into 15mls of 2xTY/20% glycerol. Single stranded DNA derived from each pool was sequenced with the cligonucleotides D1.3L40

3 and LINKSEQ1 respectively, as described above, and shown to be correctly randomised. D1.3L40:

5' CAG GAG CTG AGG AGA TTT TCC 3' 10 LINKSEO1:

5' TOC GCC TGA ACC GCC TCC ACC 3'

Preparation of rescued phage for affinity purification

10-20µl of bacteria derived from each mutagenised pool (plate scrapes) was used to inoculate 5mls of 15 TY/Glu/Amp. All bacterial growth was at 37°C. After 2-3 hours growth, 1ml was diluted in 5mls of prewarmed TY/Glu/Amp and infected by addition of 0.5 mls of a 200 fold concentrate of the M13K07gIII \(\triangle \text{No.3 preparation} \) described in example 34. After I hour of infection the 20 cultures were centrifuged at 4,000 f.p.m. for 10 minutes, resuspended in 2xTY, 100ug/ml ampicillin, incubated for a further hour, transferred to 500 mls of 2xTY medium containing 100 µg/ml ampicillin, 50 µg/ml kanamycin and 25

The remaining steps of phage grown for 16 hours. preparation were as described in example 44. Phage were finally dissolved in 10mM Tris, 1mM EDTA pH7.4 at 1/100th the original culture volume.

Affinity purification

10mls of turkey egg lysozyme at a concentration of 30 10mg/ml in 0.1M NaHCO3, 0.5MNaCl pH8.3 was mixed with an equal volume of swollen Cyanogen Bromide Activated Sepharose 4B (Pharmacia), covalently linked and washed according to manufacturers instructions. Before use this matrix (TEL-Sepharose) was washed with 100 volumes of PBS 35 followed by 10 volumes of PBSM. The TEL-Sepharose was resuspended in an equal volume of PBSM and 1ml was added

to 1ml of a 50 fold concentrate of phage in PBSM and incubated on a rotating platform for 30 minutes at room temperature. The actual phage used for this step was 40 prepared by mixing equal volumes of the independent preparations of the three randomised pools (D1.319192. D1.3HIO1 and D1.3L32). After this binding step, the

suspensions were loaded onto a disposable polypropylene column (Poly-Prep columns, Bio-Rad) and washed with 200 45 volumes of PBS containing 0.1% Tween 20. Bound phage were eluted with 1ml of 100mM triethylamine and neutralised with 0.5ml 1M Tris (pH7.4). A dilution series was prepared from the cluate and used to infect TG1 cells and plated out on TY plates containing 100µg/ml 50

ampicillin, 2% glucose. Plates carrying approximately 100 colonies were scraped into 3mls of 2mTY, 20% glycerol and stored at -70°C. 10ul of this was used to initiate a second round culture which was rescued with M13K07gIII/1 No.3 as described above (using a final culture volume of 100mls). Second and third round affinity column purification steps were carried out as described above for the first round.

Analysis by ELISA

40 colonies derived from the third round of column purification on TEL-Sepharose were picked into a 96 well plate and grown overnight at 37°C in 200µl of TY/Amp/Glu.

Phagemid particles were rescued and prepared for ELISA as described in example 18. ELISA plates were costed overnight at 37°C with hen egg lysozyme (HEL) or turkey egg lysozyme (TEL) at a concentration of 200µg/ml in 50mM NaHCO₂ pH9.6 ELISAs were carried out as described in example 18.

After 15 minutes incubation in substrate, 13 clones were found to be negative (ODc.0.05 on HEL and TEL). In all positives, a signal of 0.1-0.78 was scored on HEL with the exception of one where signal on HEL was 0.078 but signal on TEL (OD 0.159) brought it in to the positive group. The control phagemid preparation had a percentage ratio of signal TEL:HEL of 22%. Clones were deemed to have an unaltered binding if the ratio of TEL:HEL was less than 40%. 9 clones fell into this category. 18 samples were scored as having altered binding with a ratio of signal on TEL:HEL of between 40-200%.

A dilution series was made on 10 clones which were 30 analysed by ELISA in 6 of these clones the profile of binding to HEL was the same as the original clone (pCAT3SCFVD1.3) while the signal with TEL was increased (see figure 50 clone Al). In the remaining 4 clones, the increased signal with TEL was accompanied by a decrease in signal on HEL isse figure 50 clone A4).

Competition with soluble antigen All of the isolated clones retained binding to HEL to varying extents. In order to determine whether a

soluble antigen could compete with the immobilised antigen, a parallel experiment was carried out, as above, but with the addition of hen egg lysoxyme (Img/ml) to TEL-Sepharose before incubating with the phage preparation. This experiment was carried through 3 rounds of column purification and 40 colonies were

rounds of column purification and 40 colonies were picked. None of these clones bound HEL or GEL demonstrating that the soluble antigen had been successful in competing out binding to the immobilised antigen. Example 46

50 Modification of the Specificity of an antibody by Replacement of the VLK Domain by a VLK Library derived from an Unimmunised Mouse

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When an antibody specificity is isolated it will often be desirable to alter some of its properties particularly its affinity or specificity. This example demonstrates that the specificity of an antibody can be altered by use of a different VL domain derived form a repertoire of such domains. This method using display on phage would be applicable to improvement of existing monoclonal antibodies as well as antibody specificities derived using phage antibodies. This example shows that replacement of the VL domain of scFvD1.3 specific for Hen eggwhite lysozyme (HEL) with a library of VL domains allows selection of scFv fragments with bind also to More generally this Turkey eggwhite lysozyme (TEL). experimental approach shows that specificities of antibodies can be modified by replacement of a variable domain and gives a further example of the hierarchical approach to isolating antibody specificities.

The D1.3 heavy chain was amplified from an existing construct (pSWI-VHD1.3, Ward et al., 1989 supra) by PCR using the primers VHIBACK and VHIFOR, the light chain library was amplified from a CDNA library derived from the spleen of an unimmunised mouse, which was synthesized by using the MIKFONX primers 1,2,4,5 for the first strand as in example 14. The subsequent amplification was performed with the same forward primers and the VKZBACK primer. The PCR assembly of the D1.3 heavy chain with the light chain library was mediated by the signal chain

Fy linker as described in example 14.

Cloning the assembled PCR products (scFv sequences)
was done after an additional PCR step (pull-through)
using a BACk primer providing an ApaLI site and forward
primers which contained a Not 1 site as described in
example 14. ApaLI/Not 1 digested PCR fragments were
cloned into the similarly digested vector fdCAT2 as in
sample 11. Sx10³ transformations were obtained after
electroporation of the ligation reaction into MC1061
cells.
Screening of the phage library for TEL binders was

performed by penning. Polystyrene Falcon 2058 tubes were coated (16 hrs) with 2 ml of TEL-PBS (3 mg/ml) and blocked for 2 hrs with 4 ml MPBS (PBS containing 2% skimmed milk powder). Phage derived from the library (5x10¹⁰ transducing unites) in 2 ml of MPBS (2%) were incubated in these tubes for 2 hrs at room temperature.

The tubes were washed 3x with PBS, 1x with 50 mM Tris-HCl, pH 7.5, 0.5 M NaCl; 1x with 50mM Tris-HCl, pH 7.5, 0.5 M NaCl; 1x with 50mM Tris-HCl, pH 7.5, 0.5 M NaCl; 1x with 50mM Tris-HCl, pH 7.5 but 100 mM trie-thylamine. Eluced phages were eluced with 100 mM trie-thylamine. Eluced phages were taken to infect TGl cells, the cells were plated on

50 2xTY plates containing 15 µg/ml tetracycline and grown for 16h. The colonies were scraped into 25ml of 2xTy medium and the phages were recovered by PEG

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precipitation. After a second round of selection for TEL binders ELISAs were performed as described (example 2).

Analysis of 100 clones from the library before affinity selection by ELISA on plates coated with TEL showed no binders. In contrast, after two rounds of selection for TEL binding phages about 10% of the phage clones showed positive ELISA signals. ELISA signals were scored positive with values at least two fold higher than the fdCAT2 vector without insert. A more detailed analysis of binding properties of TEL binding phages is

10 shown in figure 51.

As shown in figure 51, several clones were found which bind equally to TEL and HEL in contrast to the original D1.3 scFv, which binds almost exclusively to HEL. None of the clones bound to BSA. These findings indicate that the specificity of these scrvs was broader in comparison to D1.3, since both lysozymes (HEL and TEL) are recognized, but specificity for lysozyme was retained since other BSA was not recognized. The deduced amino acid sequences (derived by DNA sequencing) of two light chains from clones MF1 and M21, which correspond to

clones 3 and 9 in figure 51 are shown in figure 52.

In the case of isolated antibodies the experimental approach as described in this study may be particularly 25 useful if recognition of a wider range of different but closely related antigens is desired. For example. monoclonal antibodies against viral antigens viral antigens like V3 loop of HIV-1 gp120 are in most cases quite specific for one particular virus isolate because 30 of the variability in this part of the HIV-1 env cene. The modification of such antibodies in the way described in this example may lead to antibodies which cross react with a wider range of HIV-1 isolates, and would

therefore be of potentially higher therapeutic or 35 diagnostic value.

A similar approach could be taken in which a light chain variable domain of desired properties is kept fixed and combined with a library of heavy chain variable domains. Some heavy chains, for example VHD1.3 retain binding activity as single domains. This may allow a strategy where VH domains are screened for binding activity when expressed on phage and then binding domains

combined with a library of VL domains for selection of

suitable light chain partners.

45 Example 47

Selection of a Phage Antibody Specificity by Binding to an Antigen attached to Magnetic Beads. Use of a Cleavable Reagent to allow elution of Bound Phage under Mild Conditions

When a phage antibody binds to its antigen with high affinity or avidity it may not be possible to elute the phage antibody from an affinity matrix with a molecule

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related to the antigen. Alternatively, there may be no suitable specific eluting molecule that can be prepared in sufficiently high concentration. In these cases it is necessary to use an elution method which is not specific to the antigen-antibody complex. Unfortunately, some of the non-specific elution methods disrupt phage structure, for instance phage viability is reduced with time at pH12 (Rossomando, E.F. and Zinder, N.D. J. Mol. Biol. 36 387-399 1968). A method was therefore devised which allows elution of bound phage antibodies under mild conditions (reduction of a dithiol group with dithiothreitol) which

do not disrupt phage structure.

Target antigen was biotinylated using a cleavable biotinylation reagent. BSA conjugated with 2-phenyl-5oxazolone (O. Makela et al. supra) was modified using a 15 biotinylation reagent with a cleavable dithiol group (sulphosuccinimidyl 2-(biotinamido) ethyl-1.3dithiopropionate from Pierce) according to the This biotinylated antigen manufacturers instructions. was bound to streptavidin coated magnetic beads and the 20 complex used to bind phage. Streptavidin coated magnetic beads (Dynal) were precoated with antigen by mixing 650µg of biotinylated OX-BSA in 1 ml PBS, with 200µl of beads Free antigen for at least 1 hour at room temperature. was removed by washing in PBS. One fortieth of the 25 complex (equivalent to 5µl of beads and an input of 17.5 ug of OX-BSA) was added to 0.5ml of phage in PBSM (PBS containing 2% skimmed milk powder) containing 1.9x1010 phage particles mixed at the ratios of pAbD1.3 directed against lysozyme (example 2) to pAbNQ11 directed against 30 2-phenyl-5-oxazolone (example 11) shown in Table 12.

After 1 hour of incubation with mixing at room temperature, magnetic beads were recovered using a Dynal MPC-E magnetic desperation device. They were then washed in PBS containing 0.5% Tween 20, (3x10 minutes, 2x1 hour, 2x 10 minutes) and phage eluted by 5 minutes incubation in 50µl PBS containing 10mM dithiothreitol. The eluter was used to infect TG1 cells and the resulting colonies

probed with the oligo NQ11CDR3
40 (5' AAACCAGGCCCCGTAATCATAGCC 3')

derived from CDR3 of the NQ11 antibody (This hybridises to pabNO11 but not pab D1.3).

A 670 fold enrichment of pAbNQ11 (table 12) was achieved form a background of pAbD1.3 in a single round of purification using the equivalent of 17.5µg of biotinylated OX-BSA.

This elution procedure is just one example of an elution procedure under mild conditions. A particularly advantageous method would be to introduce a nucleotide sequence encoding amino acids constituting a recognition site for cleavage by a highly specific protease between the foreign gene inserted, in this instance a gene for an

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antibody fragment, and the sequence of the remainder of gene III. Examples of such highly specific proteases are Factor X and thrombin. After binding of the phage to an affinity matrix and elution to remove non-specific binding phage and weak binding phage, the strongly bound phage would be removed by washing the column with protease under conditions suitable for digestion at the cleavage site. This would cleave the antibody fragment from the phage particle eluting the phage. These phage would be expected to be infective since the only protease 1.0 site should be the one specifically introduced. Strongly binding phage could then be recovered by infecting e.g.

E.coli TG1 cells.

Example 48

15 Use of Cell Selection to provide an Enriched Pool of Antigen Specific Antibody Genes, Application to reducing the Complexity of Repertoires of Antibody Fragment displayed on the Surface of Bacteriophage
There are approximately 10¹⁴ different combinations

of heavy and light chains derived from the spleen of an immunised mouse. If the random combinatorial approach is used to clone heavy and light chain fragments into a single vector to display scFv, Fv or Fab fragments on phage, it is not a practical proposition to display all 25 1014 combinations. One approach, described in this example, to reducing the complexity is to clone genes

only from antigen selected cells. (An alternative approach, which copes with the complexity is the dual combinatorial library described in example 26).

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The immune system uses the binding of antigen by surface immunoglobulin to select the population of cells that respond to produce specific antibody. This approach of selecting antigen binding cells has been investigated to reduce the number of combinatorial possibilities and so increase the chance of recovering the original combination of heavy and light chains.

The immunological response to the hapten 4-hydroxy-3-nitrophenylacetic acid (NP) has been extensively studied. Since the primary immune response to NP uses 40 only a single light chain the applicants were able to examine the use of the combinatorial method using a fixed light chain and a library of heavy chains to examine the frequencies genes that code for antibodies binding to NIP

(4-hydroxy-3-iodo-5-nitrophenylacetic acid). 45 applicants have thus used this system to investigate the merits of selecting cell populations prior to making combinatorial libraries for display on phage.

Methods

2.1 Hapten conjugates

Chick gamma globulin (CGG, Sigma, Pople, UK) and Bovine serum albumen (BSA, Boehringer, Mannheim, Germany) were conjugated with NP-O-succinimide or NIP-caproate-O-

succinimide (Cambridge Research Biochemicals, Northwich, UK) based on the method described by Brownstone (Brownstone, A., Mitchison, N.A. and Pitt-Rivers, R., Immunology 1966. 10: 465-492). The activated compounds were dissolved in dimethylformanide and added to proteins in 0.2 M sodium hydrogen carbonate. They were mixed with constant agitation for 16 hours at 4°C and then dialysed against several changes of 0.2 M sodium hydrogen carbonate. They were finally dialysed into phosphate buffered saline (PBS). The conjugates made were NP12CGG, NIF10BSA. The NIF10BSA derivative was subsequently biotinvlated using a biotinylation kit purchased from

2.2 Animals and immunisation

Mice of the strain C57BL/6 were immunised by intraperitoneal injection of 100pg NP-CGG in Complete

Amersham (Amersham International, Amersham, UK).

Freunds Adjuvant at 10 weeks of age.

2.3 Spleen preparation

Seven days after immunization cells from the spleen
were prepared as described by Galfre and Milstein
(Galfre, G. and Milstein, C. Methods Enzymol. 1981. 73:346). Red cells were lysed with ammonium chloride (Boyle,
W. Transplantation 1968.6:71) and when cell selection was
performed dead cells were removed by the method described
by von Boehmer and Shortman (von Boehmer, H. and
Shortman, K, J. Immunol, Methods 1973:1:273). The cells
were suspended in phosphage buffered saline (PBS), 1%

were suspended in phosphage buffered saline (PBS), 1% Bovine serum albumen, 0.01% sodium azide; throughout all cell selection procedures the cells were kept at 4°C in this medium.

2.4 Cell Solution

Biotinylated NIF-BSA was coupled to streptavidin coupled magnetic beads (Dynabeads M280 Streptavidin, Dynal, Osio, Norway) by incubating 10⁸ beads with 100µg of biotinylated protein for 1 hour, with occasional agitation, and then washing five times to remove unbound antigen. The coupled beads were stored at 4°C in medium until required. For selection of antigen binding cells the cells (2-4x10'mm) were first incubated for 30 minutes with uncoupled beads, at a bead: cell ratio of 1:1, to examine the degree of non-specific binding. The beads were then separated by placing the tube in a magnetic device (MPC-E Dynal) for 3-5 minutes. The unbound cells were removed and then incubated with NIP-

45 BSA coupled magnetic beads, at a bead:cell ratio of 0.1:1, for 60 minutes, with occasional agitation. The beads and rosetted cells were separated as described above. The beads were then resuspended in 1 ml of medium and the separation repeated; this process was repeated 5- 7 times until no unbound cells could be detected when

counted on a haemocytometer.

For the depletion of surface immunoglobulin positive

cells the cells were incubated with 20µg biotinylated goat anti-mouse polyvalent immunoglobulin (Sigma, Poole, UK). The cells were then washed twice with medium and added to streptavidivin coupled magnetic beads at a beed to cell ratio of 30:1. After 30 minutes incubation the beads and rosetted cells were separated by applying the magnetic device three times - taking the supernatant each time.

2.4 DNA/CDNA preparation, FCR amplification and cloning
DNA was prepared by a simple proteinase-K digest
method that was particularly convenient for small numbers
of cells (FCR Protocols: A Guide to Methods and
Applications. Ed Innis M.A., Gelfand D. H., Sninsky J.J.

and White T. J. Academic Press). RNA preparation and 15 subsequent cDNA synthesis was performed as described by Gherardi et al (Gherardi E., Pannell R. and Milstein C. J. Immunol. Methods, 1990. 126:61-68). FCR and cloning of the heavy chain libraries was performed using the primers and conditions described by Ward et al (Ward,

20 E.S., Güssow, D., Griffiths, A.D., Jones, P.T. and Winter, G., Nature, 1989. 341: 544-546): 40 cycles of PCR amplification were performed. The VH and Fv expression vectors used were adapted from those previously described by Ward et al. They were both

25 subcloned into pUC119 (Veira and Messing see later) and the Fv expression vector was modified to include a germline lambda-1 light chain (obtained as a gift from T. Simon (originally cloned by Siegfried Weiss, Basel Institute of Immunology)). The vector is shown in Figure 30 53.

2.5 Expression and ELISA

For screening single colonies were picked into individual wells of microtitre plates (Bibby) in 200ml 2 * TY/Ampicillin 100µg/ml/0.1% glucose and then incubated at 37°C for 5-6 hours with agitation, Isopropyl-β-D-35 thiogalactopyranoside (IPTG, Sigma, Poole, UK) was then added to a final concentration of 1 mM and the incubation continued for a further 16 hours at 30°C before harvesting the supernatants. The wells of Falcon ELISA 40 plates (Becton Dickenson, N.J., USA) were coated overnight at room temperature with NIP10-BSA (40µg/ml in PBS) and then blocked with 2% skimmed milk powder in PBS for 2 hours at room temperature. The bacterial supernatants were added and incubated at room temperature 45 for 1 hour and then the plates were washed three times with PBS. Peroxidase conjugated-Goat anti-mouse lambda-

chain (Southern Biotechnology, Birmingham, USA) was added and again incubated for 1 hour at room temperature before washing six times with PBS and then developing with 2.2'-50 Azino-bis (3-ethylbenzthiazoline-6-sulfonic acid) (Sigma, Poole, UK) as the peroxidase substrate. The optical density at 405nm was measured using a Thermomax

PCT/GB91/01134

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microplate reader (Molecular Devices, Menlo Park, USA) after 30 minutes. Western blotting using the C-terminal myc tag as described in example 27.

3.1 Comparison of RNA/DNA and antigen selected cells

- The results of antigen selection are shown in Table Less than 1% of cells bind to NIP-BSA coated beads and the non-specific binding is very low. Assessment of the proportion of expressed genes from each VH library using western blotting showed that full length VH domains were expressed in 95% (19/20) of all clones when RNA was 10 used as the starting material but only 60% (12/20) of clones when DNA (either selected cells or from total spleen) was used as the starting material. difference probably results from the fact that many rearranged pseudogenes could be amplified with our primers
- 15 and it appears that there must be some degree of selection, at the level of transcription, for functional denes.
- A variable number of clones from each type of library were screened for the production of Fv fragments 20 that bound to NIP. Initial screening ELISAs were performed and positives taken to include those with an optical density of at least twice the background. initial positives were retransformed and the binding
- checked in duplicate; it was confirmed that the binding 25 was specific to NIP and not to BSA. The frequency of confirmed positive NIP binding clones for each starting material are shown in Table 14. Using DNA as the starting material for the PCR amplification is
- approximately equivalent to sampling the cells present as 30 there is only one functional re-arranged heavy chain gene and at most one re-arranged pseudogene per B-cell. Amplifying from the RNA of an animal of course biases the reperroire to the reacting B-cells and in a recently
- immunised animal this would be expected to give some bias 35 The data in Table 14 clearly towards the immunocen. shows how powerful this selection is with the number of anticen specific genes being enriched at least 96 fold when RNA made one week after primary immunisation is used as the starting material. The data also show that 40
- selection for antigen binding cells also provides an alternative powerful method of selection for the required genetic starting material.
 - 3.2 Comparison of Total Spleen/surface immunoglobulin depleted Spleen To examine the cellular basis of the selection
- achieved by using RNA as the starting material we depleted the spleen of surface immunoglobulin positive cells using biotinylated anti-polyvalent immunoglobulin and streptavidin conjugated magnetic beads. Prior FACS analysis had demonstrated that this method removed over 96% of surface immunoglobulin positive cells. RNA was

prepared from both surface immunoglobulin depleted and non-depleted factions of a spleen and VH libraries made from each. The ELISA results (Table 14) show that the number of positives is certainly not decreased by this depletion suggesting that the major portion of the selective effect of using RNA may come from surface immunoglobulin negative G-cells (probably plasma cells). Conclusions

The applicants have demonstrated the importance of the amplification of specific RNA produced by immunisation to enable binding activity to be obtained with any reasonable frequency from a combinatorial library. The applicants have also demonstrated an alternative strategy which mimics that of the immune system itself. Using a simple method of selecting for antigen binding cells gave comparable enrichment and has the added advantage of using a broader range of genes. At first sight the random combinatorial approach would appear unlikely to produce the original combination of

20 heavy and light chain because of the vast diversity of the immunoglobulin genes. The applicants show here, however, that following immunisation, with a good antigen, 10% of the VH genes from total splenic RNA isolated come from antigen specific cells so the

25 effective size of the repertoire is greatly reduced.

This together with the fact that promiscuity of the heavy and light chains occurs (examples 21 and 22) accounts for the fact that combinatorial system does produce antigen binding clones with reasonable frequency. The data also suggests that the built of the articen precision produces.

suggests that the bulk of the antigen specific RNA comes from surface immunoglobulin negative cells which are most likely plasma cells.

The data also show that this simple method of antigen selection may be useful in reducing the complexity of the combinatorial library. In this case an enrichment of antigen specific genes of at least 55 fold has been achieved which in the normal case where heavy and light chains are unknown would result in a reduction of the complexity of the combinatorial library by a

40 factor of over 3000. A further advantage of using antigen selected cells (and amplifying from DNA to reduce any bias due to the state of the cell) is that this results in a broader range of antibody genes amplified. It may be that a simple cell selection such as that the

45 applicants have described here in combination with phage selection would be ideal. From this example it can be seen that by combining cell and phage selection methods one could reasonably expect to screen all the combinations of heavy and light chain (approximately 4x10¹⁰) and would thus be able to screen all binding

50 4x10¹⁰) and would thus be able to screen all binding combinations although this would not, at present, be possible from whole spleen (approximately 4x10¹⁴ combinations, assuming 50% B-cells).

Table 1. Enrichment of pAb (D1.3) from vector population

INPUT RATIO*	OUTPUT RATIO		ENRICHMENT ^d	
	oligob	ELISA		
pAb:fd-CAT1	pAb:total phage	pAb:total phage		
Single Round	ő .			
1:4x103	43/124		1.3×103	
1:4×10*	2/82		1.0×10°	
Two Rounds				
1:4x104	197/372		2.1x104	
1:4×10 ⁵	90/356	3/24	1.0x10 ⁵	
1:4x10 ⁶	27/183	5/26	5.9x10 ⁵	
1:4x107	13/278		1.8x10°	

Footnotes: ^aApproximately 10^{12} phage with the stated ratio of pAb (D1.3): FDTPs/Bs were applied to 1 ml lysozyme-sepharose columns, washed and eluted. ^bTG1 cells were infected with the eluted specific binding phage and plated onto TY-tet plates. After overnight incubation at $30-37^{\circ}\mathrm{C}$, the plates were analysed by hybridisation to the $^{32}\mathrm{p}$, labelled oligonucleotide VHIFOR (Ward et al op cit) which is specific to pAb D1.3. ^C Single colonies from overnight plates were grown, phage purified, and tested for lysozyme binding. ^dEnrichment was calculated from the oligonucleotide probing data.

Table 2 Enrichment of pAb (D1.3) from mixed pAb population

Input Raticl (pAbD1.3:pAbNQ11)	Output Ratio ² (pAb D1.3:Total phage)	Enrichment
Single Round		
1 : 2.5 x 10*	18/460	0.98 x 10 ³
1 : 2.5 x 10 ⁵	3/770	0.97 x 103
1 : 2.5 x 10 ⁶	0/112	**
pAb NQ11 only	0/460	~
Second Round		
1:2.5 x 104	119/170	1.75 × 104
1 : 2.5 x 10 ⁵	101/130	1.95 × 105
1 : 2.5 x 10°	102/204	1.25 × 106
1 : 2.5 x 10 ⁷	0/274	100
1 : 2.5 x 10°	0/209	**
pAb NQ11 only	0/170	**

Notes

- 1. 10^{10} phage applied to a lysozyme column as in table 1.
- Plating of cells and probing with oligonucleotide as in table 1, except the oligonucleotide was D1.3CDR3A.

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Table 3: Enzyme activity of phage-enzyme

Input	ng of enzyme or No. of phage	Rate (OD/hr)	No. of molecules of Enzyme equivalent (x10 ⁻¹¹)
Pure Enzyme	335	34	24.5
Pure Enzyme	177.5	17.4	12.25
Pure Enzyme	88.7	8.7	6.125
Pure Enzyme	44.4	4.12	.3.06
Pure Enzyme	22.2	1.8	1.5
Pure Enzyme	11.1	0.86	0.76
No Enzyme	0	0.005	0
fd-phoAm166/TG1	1.83×10 ¹¹	5.82	4.2
fd-CAT2/TG1	1.0x10 ¹²	0.155	0.112
fd-phoAlal66/KS272	7.1x1010	10.32	7.35
fd-CAT2/KS272	8.2x10 ¹²	0.038	0.027

Table 4. Affinity selection of hapten-hinding phage.

Chares binding to plath.

Random Contibuatorial Libraries (9568 (0%) 48/276 (13%) 175/188 (93%) PhDe-immunised mice (9568 (0%) 48/276 (13%) (93%) Hierarchical Libraries (979) 34/380 (92%) VII-14 / Vs. rrp Boary (979) (3%) 23/380 (7%)		\$\$156.15 \$25.56.15		Mey first round		After second round	Me
ites 6/190 (3%) 348/380 (92%) 6/190 (0%) 23/380(7%)		Random Combinatorial 13 pbDe-immunised mice Unimmunised mice	Bruries (0%)	48/376 (13%)	(75/188 (93%)	TO THE PROPERTY OF STATE OF ST	
(%0) (0.17)	1	Merarchical Edicaries VII-R/Vs. 173 Brany	64190 (3%)	3.187.380 (92%)	e destruction and a constant of the state of	or annual constant constant of many (or	# 1 form more a part amades
		VII.rep / Vr4 libary	0/190 (0%)	23/380 (7%)	,		
		Mixing of chars	88/1896 (4.6%)	5505 (57.9%)	1152/1156 (90,7%)	1296/1209 (99.8%)	9.8%)

f in panel C, manhers refer to VH-B/VK-d colonies.

* Nambers after three reinfections and cycles of growth. This control, custiting the cohann steps, continus that a spurious geowife or interityly abanture was not responsible for the enticlinean for clan. VII Weself.

Table 5

Soluble chain(a)*	bouvy chain Bouvy chain	300 Apr.	heavy chain heavy chain held than light chain
Chain as Sotuble gene III jusing ⁹ chain(s) ⁸	safv hgat chain be sch'y light chain	76 22	light chain light chain light chain light chain
2			2
2 2 3 3 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4	Signal Si		
2	222		5656
	6 6 2 7 2 - 1 1 6 6 7 2 - 1 1	pushi i (m2151) pushi ii (m2151)	66 CAP2-111 66 CAP2-18 616N III (10215) PHEN III (10215) PHEN IV (10215)

accordance with hinding data, § Reanh continued experimentally by Western blut (for Fab, see * Phage were considered to be binding if OD405 of sample was at least 10 fold greater than E.cofi 1102151 is specifically indicated, # Information deduced from genetic structure and in background in ELLSA; † E.coli Wil was used for the growth of the phage unless the use of Overview of phOx-BSA ELISA results of phage and phagemid constructions. **Евнс** 29.

	Soluble (Data from et al 1988)	enzyme Chaidaroglu	Phage (Data from	enzyme this study)
	phoArg166	phoAla166	phoArg166	phoAla166
$K_m\ (\mu M)$	12.7	1620	73	1070
Relative K_{m}	1	127	1	14.6
Relative kcat	¥.	0.397	1	0.360
Relative kcat/Km	1	0.0032	1	0.024

Table 6. Kinetic parameters of soluble and phage-bound alkaline phosphatase. Relative values of $k_{\rm cat}$ and $K_{\rm m}$ for the soluble enzyme and for the phage enzyme were derived by comparing with the values for wild type enzyme (phoArg166) and the phage-wild type enzyme (fdphoArg166).

Enzyme Activity of Phage Samples

SAMPLE (Construct:host)	INPUT PHAGE PARTICLE (pmol:	RATE (pmol substrate converted/min)	SPECIFIC ACTIVITY (mol substrate converted/mol phage/min)
fdphoArg166 :TG1	2.3	8695	3700
fdphoAlal66 :TGl	5.6	2111	380
fdphoAla166 :KS272	1.8	2505	1400
fdCAT2: TG1	3.3	<1	<0.3
fdCAT2: KS272	5.6	70	12

Table 7

Table 8. Affinity chromatography of phage-enzymes

SAMPLE	INFECTIVITY (Percentage of phage particles which are infectious)	INPUT PHAGE PARTICLE (×10 ⁹)	OUTPUT PHAGE PARTICLE (x10°)
fdphoArg166	0.37%	5160	30
fdphoAla166	0.26%	3040	90
fdCAT2	4.75%	4000	2

Nucleotide mutation	Amino acid mutation	Number
(base position) 308	Ala->Val (VH FR3)	3
703	Tyr->Asp (VL CDR3)	1
706	Ser-> Gly (VL CDR3)	1
724	Gly-> Ser (VL FR4)	21
725	Gly-> Asp (VL FR4)	3
734	Thr-> Ile (VL FR4)	

Table 9 Mutations in scFvB18 selected by display on phage following growth in mutator strains

Table 10(1)Oligonuclatide primers used for PCR of human immunoglobulin genes

Human VII I	Oligo Name
Back	
Primers	
	Sequence

HuVIIIaBACKSti HuVH2aBACKSti HuVH3aBACKSti HuVH4aBACKSti HuVH5aBACKSti	HaVIIIaBACK HaVH2aBACK HaVH3aBACK HaVH3aBACK HaVH4aBACK HaVH5aBACK
51-GITC CITC GCA ACT GCG GCC CAG GCG GCC AVIG GCC CAG GTB CAG CTB GTB CAG TCB (32-3) 51-GITC CITC GCA ACT GCG GCC CAG GCG GCC AVIG GCC CAG GTB CAG CTB AGG GAG TCT G3-3) 51-GITC CITC GCA ACT GCG GCC CAG GCG GCC AVIG GCC CAG GTB CAG CTB GTB GAG TCT G3-3) 51-GITC CITC GCA ACT GCG GCC CAG GCG GCC AVIG GCC CAG GTB CAG CTB TCB CAG GTC TCB C3-3) 51-GITC CITC GCA ACT GCG GCC CAG GCC AVIG GCC CAG GTB CAG CTB TCB CAG TCB TCB C3-3) 51-GITC CITC GCA ACT GCG GCC CAG GCC AVIG GCC CAG GTB CAG CTB TCB CAG TCB TCB (32-3) 51-GITC CITC GCA ACT GCG GCC CAG GCC AVIG GCC CAG GTB CAG CTB TCB CAG TCB TCB (32-3)	5'-CAG GIG CAG CIG GIG CAG ITCT GG-3' 5'-CAG GIG CAG CIG GIG GAG ITCT GG-3' 5'-CAG GIG CAG CIG GAG TCT GG-3' 5'-CAG GIG CAG CIG CAG GAG TCT GG-3' 5'-CAG GIG CAG CIG CAG CAG TCT GG-3'

Human JH Forward Primers

HuJH6FOR	HuJH4-5FOR	HuJH3FOR	HuJHI1-2FOR
5'-TGA	5'-TCA	5'-ICA	5'-TG
8	8	ACA	9
88	38	300	0,00
GGT	8	63	GT
98	360	R	000
8	000	CAT	C
8	8	TOT	037
OX-31	TC-3	000-3	000-31

Human Heavy Chain Constant Region Primers

Table 10 (11"

Human Jk Forward Primers

HulgG1-4CHFOR HulgMFOR Human Vk Back Primers	5'GIC	88	CAC CIT GGI GIT GCI GGG CIT-3'	38	GIT GIT	98	7 8	98
HuvkiaBACK	51-020	Alic	S	NG	Š	CK CK	TCT 02-3	Ŗ
HuVkiaBACK HuVk2aBACK	21-030	GIT	08 08 08 08			0 0 0 0	85.5	११
HuVĸlaBACK HuVĸ2aBACK HuVĸ3aBACK	5'-GAC AIC CAG 5'-GAI GIT GIG	A A A	2 2 2			888	10 10 10 10 10 10 10 10 10 10 10 10 10 1	នុនុនុ
HuVkiaBACK HuVk2aBACK HuVk3aBACK HuVk4aBACK	51-026 51-026		9998			8888	西西西西	8888
HuVklaBACK HuVk2aBACK HuVk3aBACK HuVk4aBACK HuVk4aBACK	51-030 51-030 51-030 51-030					88888	拉拉拉拉拉	88888

Hujk2BACKNot Hujk4BACKNot Hujk4BACKNot	IIIujkiBACKNot	HuJk5FOR	Hujk4FOR	Hujk3FOR	HuJk2FOR	HujxIFOR
5'-GAS TOA FIC TICS ACT TICS CCC CCC ACC FIT (AVI CTC CAC CIT GAI CXC-3' 5'-GAS ICA FIC TICS ACT TICS CCC CCC ACG FIT (AVI CTC CAC CIT GAI CXC-3' 5'-GAS TOA FIC TICS ACT TICS CCC CCC ACG FIT (AVI CTC CAC CIT GAI CXC-3' 5'-GAS TOA FIC TICS ACT TICS CCC CCC ACG FIT AVI CTC CAG TICS TICT CXC-3'	TOA THE THE ACT THE GOD ONE ACT THE WAT THE COSE CHI	5'-AG TIT AAT CTC CAG TCG TGT CCC-3'	51-ACG TIT CAT CIC CAC CIT GGT CUC-31	5'-ACC TITE GAT AIC CAC TITE GGT CCC-3'	5'-ACC THE GAT CIC CAS CIT GGT CCC-3'	5'-ACC TIT GAT THE CAS CIT GGT CCC-3'

Human K Constant Region Primers

Table 10 (1111

ST-MA CRUTEC CLY GITHAN GLY CIT-31 I ST-GROTEN THO THE ACT THE CASO CAST HAN THE AND CHUTEC OF GITHAN GLY CYT-31 ST-GROTEN THO THE ACT THE CASO CAST AND CHUTEN COFF GITHAN ACT CYT-31	HaCkfOKNot2	HuCkFORNott	HUCKPOR
	A THE PART AND MEN AND ARE AND AND AND ARE AND THE WASHINGTON WHOLE	5'-FR WALLE WE	STARR CRUBE COS GENERA GOT CREST

Human & Back primers

Hu\(\delta\)6BACK	Huλ5BACK	HaA4BACK	Пидзываск	Πuλ3aBACK	Huλ2BACK	HnλIBACK
23-VVI.	51-CM3	51-030	5'-ICT	5'-ICC	51-076	51-036
775	83	CIT	TOP	TAT	TCT	T
AIG	GEG	ATA	000	GIG	88	9
CIG	CIC	CIG	SE	CIG	CIG	TTC
LW	ACT	ACT	ACT.		MCT	200
30	35	CAA	CPG CPG	S	0.50	080
22	8	8	8	8	S	200
CCC CA-3'	TC-31	CCG CC-3*	GBC CC-3*	CCA CC-3*	OCT 90-31	OUG CC-3'

Human & Forward Primers

Hu JAHORNOT Hu JA2-3FORNOT Hu JA4-5FORNOT	Hu BA-SFOR Hu BA-SFOR
5"-GMG TEA THE THE ALT THE GREE COE ACT THE FRE GREETH GREETH GREETH CLEFF. 5"-GMG TEA THE THE ALT THE GREE COE ACT THE FRE GREETH GREETH CLEFF. 5"-GMG TEA THE THE ALT THE GREE COE ACT ACT THA FRE GREETH GREETH GREETH CLEFF.	51-ACC TRG GAC GOT CAY: CYF GOT COC-31 51-ACC TRG GAC GOT CAY: CYF GOT COC-31 51-ACC TRA ANC GOT GOT CYC GOT COC-31

Human & Constant Region Primers

Table is

Reverse IgC1-4C111 primer for Fab linker RHuji 16 RHuJH4-5 KH IuJH 13 RHIJHI-2

5'-GEN OWN OSS S'-GAA COC TOG TCA 51-COA CAN TICK TOA 5'-GCA COC TIGG TICA S 8 8 8 101 SS IG TCF ij 8 8 CIT 8 CAR CIRC C-3' 9 8

GIG G-3 GTG G-31

Reverse Vk for sch'v linker

RHINGCH-4CHHOR 51-ANG CLE NYC AND NYC AND GIRG GAC-31

RHuVk6aBACKFv RHuVk5aBACKFv RIIuVK4aBACKFv RHuVkJaBACKFv RIIuVr2aBACKFv RHuVklaBACKFv 51-034 SI-CH CHO 5'-GEN GIC 21-CH SI-OH 57-GCA CAC TOS 000 8 8 TGA 18 S 13 TCA GIC NGC GIG NGT CIC SIC OIC. SIC AIC PAC AIC TGG ATC MA 9 Š ACA CA Š AIG ATT T AGA ATG ATT Ŕ ğ 301 ä 300 Ŕ CAT CUT CUT-3' CPAT CALL COR LANGE CAR CXX CX-31 GI 91 8 COS (C-3* 003 00-3 8

Reverse Vk for Fab linker

RHuVk6aBACKFab RITh V KSaBACK Fab RHuVk4aBACKFab RI Iu V KJaB A CK Fab RI In Vx2aBACK Fab RHuVxlaBACKFab 2,703 21-32 CKC WYY.S 5'-03 57-QM W.S.-S 80 25 CAK. CHU THE CHU MIC GIC TOS GIC AIC W. 13 XX 3 380 CTC CRC GE JEN ! 383 200 MC AXA CIC. N.K. N. N. MA 103 GIT MY AIG MI ACA MC TOTAL OF CALL TES GY: ATT GT GE3 TO TOUGHT AND AND BOTH 100 103 GT 98 38 W. 03. 653 18-48 JULY 1818 ATT GT GE3

Reverse VA for svFv linker

RIJuV\BACK3bFv RHnVlBACK6Fv RIInVABACK5Fv KHuV\BACK4Fv RHuVABACK3aFv RIIuV\BACK2I'v RITUVABACKIFV 51-66 280-15 25-033 IES-1S 51-CA 51-030 93 8 300 8 8 33 ORD WEST 303 100 S ŢÇ, 103 8 OEC. 900 SIC 910 CIC 350 Ã H 8 XX AGA ACC ACA AS. ATA ANA THE CAST CHELLES AND CHELLES CAND-31 ATA K.N. ACA 93 A. W. 9 200 B GYA. 78 8 TOC GAT 18 8 000 S ğ 2 CAT CAT CAT CLE US COR US CON COR-3. COS OTA CUE CUA AND SEE AND SEE CES CEN CC CC 2000 CLIS CLE 08 CCC CLY CAC-3. CA CNI-3 9G-3 GAR-34 (A)(2-3

Reverse VA for Fab linker

RI In VABACK4Fab RHuVABACK3bFab KHuVABACK3aFab RHuV\BACK6Fab RHuV\BACK5Fab RHuV\LBACK2Fab RHuVABACKIFab 51-60 51-05 5.-GI 5'-OJ 51-030 STATE OF THE OTHER THE THE WAS AND THE OTHER OF THE STATE S 5,788 98 8 CC 8 8 8 E.M. TON (E) G 8 TY YE STO CHC. GIC SIC 900 GE K 30 Š AGA GA AAC NO: NO ATA TCA 8 ACA 30 R MG S Sel 9 9 BOL 3 8 98 103 103 330 88 8 8 8 OCC AND OLD ATC AIK MIC Et Li MY SE OF GI 1,50 3.4 23 (37T E.O. 33 9 TOS 077-3 TFF 0.7-3 "E-4"D SEE 17 (3 T)-3" 103 (3.74-3" TGG GCA-3*

Table II. Deduced protein sequences of heavy and light chains selected from unimmunized library

Oxazolone binder

PICHL CBTH VHIS. 4 CVOLVQCCDASVKERGASVKVSCKASSYTTET HEAVY CHAIN SICIS SMINE WYCHLFOIRPELLIK BEHRRIS GIFTRESSESSISTERILGITERCHERKARYC CAMBIE CONTROL NAMED AND STATE OF THE PROPERTY. WISHINGSTEXACKING RETWISTSTEETSTWYSSLEELESKYTSVYTSVYTCVR HIMINGALT. ASSESSAGIAA

CIVEXIE

85A Binders

8.8.38 FIGHT CHYINS HEAVY CHAINS 43,8

ORCIVOSCINANDECENTRICENSPRINTES SACRE ENACHER AND AND SERVICE STREET

SHERI/TODPAUBURLOQPURITE SAXXENTECES

XINTAMODENDOME

CKANKES

GIPDREGGSSSGHASLTINGAGGSGSGADYYC

HERDSSCREE

CCSAAA

RPTISKEREETTIMINGORGIRAETTAVYYCAK

A CASSSAN

22XXXXXXX

BELLYRIC AAVER OMOTIVOSTRIANKUSOOSTINISCUODARES

OMOTIVOSCULENKUSOOSTINISCUODARES

OMOTIVOSCULENKUSOOSTINISCUODARES STREAMSTRIB SUCKE SAMAN SAMAN SAMAN

152

Lysozyme binders:

T. BIRA VELO. 1

1. \$17A T.OIMA

OSALITOPAGUSGSFOQSITISC DELLISCOSCOSCOSCOSCILIAN SHELTODPARSVAPOUTVRITO SALLANDANGTOS GOLIALS

AITTHAYSOIRBOLM BALGARDIGESEES SITTHAYSOIRGAM RAALACCAGLESSEL

SALKSBATEGED

AIATHAVOOANDOAN

CHARRES BAMMAB SALIMAN

SHINKE CHAINS VH16.1 VH13.1

WILMSTEDGER ENGINE TOND SINCE SANDERS ENWERTEN STATE OF THE PARTY OF A I'T DK A WHO CHOOLE MASTICS

MINCEPEDROLLEMIG DIMENSACRATION OF LIPPIXXXXXXXXXXXXXX EXBERREASICEMENTS IN SCHESKACLENBELL SYTHREALTAINED

CALIENTERITENTERITENTERITENE CALIENTERITERITENTERI RVTMEVDTSKN/JSSL/RLKSVTAADIBAN/PCAR

AWAILDONY

ACABASHBAB MEMACHALA BEARTSHALLSCHOOL

COSTA PENALTY

1.3.5% TESTABLE KISKUSKOTKI.

GISHKESKSKSKTREITTISELQPKERKKYYC CANDERS SERVING LUISE LA SERVING SERVI CALCAL CONTROL INTERCORPORATION CO.

90XXX

Table 12

Enrichment of pAbNQ11 from pAbD1.3 background by affinity selection using Ox-BSA biotinylated with a cleavable reagent and binding to streptavidin magnetic heads

Input Ratio I (pAbD1.3:pAbNQ11)	Output Ratio ² (pAh NQII: Total phage)	Enrichment
******	61/197	690
22350:1	5/202	544

 ^{1. 1.9}x10¹⁰ phage in 0.5ml mixed for lhour with 5µl streptavidin-magnetic heads precoated with antigen (OX-BSA).
 2. Colonies prohed with the oligonucleotide NQ11CDR3

13 Table: Results of antigenic cell selection

Table : Results of Fv NIP binding ELISAs from selected cell populations:

	Positives	*Degree of
		Enrichment
Cell Population		
DNA from total spicen	0/940	v 4
RNA from total Spleen	29/282	> 96
DNA from antigen binding ceils	17/282	>56
Surface Ig Selection		
RNA fromSurface Ig	8/94	~

4/94

RNA from total Spiecn

^{*} Degree of enrichment compared to total DNA.

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CLAIMS

A method of producing a multimeric member of a specific binding pair (sbp), which method comprises: expressingt in a recombinant host organism a first polypeptide chain of said sbp member or a genetically diverse population of that type of sbp member fused to a component of a secreted replicable genetic display package (radp) which thereby displays said polypeptide at the surface of the package, and expressing in a recombinant host organism a second polypeptide chain of said multimer and causing or allowing the polypeptide chains to come together to form said multimer as part of said rgdp, at least one of said polypeptide chains being expressed from nucleic acid that is capable of being packaged using said

- component therefor, whereby the genetic material of each A method according to claim 1 wherein both said chains are expressed in the same host organism.
- A method according to claim 2 wherein said first and second chains of said multimer are expressed as separate chains from a single vector containing their respective nucleic acid.

said rgdp encodes a said polypeptide chain.

- A method according to any one of claims 1, 2 and 3 wherein at least one of said polypeptide chains is expressed from a phage vector.
- A method according to any one of claims 1 to 4 wherein 30 S. at least one of said polypeptide chains is expressed from a phagemid vector, the method including using a helper phage, or a plasmid expressing complementing phage genes, to help package said phagemid genome, and said component 35 of the rgdp is a capsid protein therefor.
 - A method according to claim 5 wherein said capsid protein is absent, defective or conditionally defective in the helper phage.

A method according to any one of the preceding claims

- which comprises introducing a vector capable of expressing said first polypeptide chain into a host organism which expresses said second polypeptide chain in free form, or introducing a vector capable of expressing said second 45 polypeptide in free form into a host organism which expresses said first polypeptide chain.
- A method according to any one the preceding claims 50 wherein each said polypeptide chain is expressed from nucleic acid which is capable of being packaged as a rgdp using said component fusion product, whereby encoding nucleic acid for both said polypeptide chains are packaged in respective rgdps.

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9. A method according to any one of the preceding claims wherein the nucleic acid encoding at least one of said first and second polypeptide chains is obtained from a library of nucleic acid including nucleic acid encoding said chain or a population of variants of said chain.

10. A method according to claim 9 wherein both the first and second polypeptide chains are obtained from respective said libraries of nucleic acid.

11. A method of producing a member of a specific binding pair (sbp) from a nucleic acid library including nucleic acid encoding said sbp member or a genetically diverse population of that type of sbp member, which method comprises:

expressing in recombinant host cells polypeptides encoded by said library nucleic acid fused to a component of a secreted replicable genetic display package (rgdp) or in free form for association with a polypeptide component of said sbp member which is expressed as a fusion to said rgdp component, so that the rgdp displays said sbp member in functional form at the surface of the package, said library nucleic acid being contained within the host cells in a form that is capable of being packaged using said rgdp component, whereby the genetic material of an rgdp displaying an sbp member contains nucleic acid encoding said sbp member or a polypeptide component thereof.

12. A method of producing a member of a specific binding pair (sbp), which method comprises:

expressing in recombinant host cells nucleic acid encoding said sbp member or a genetically diverse population of that type of sbp member, wherein the or each said sbp member or a polypeptide component thereof is expressed as a fusion with a component of a secreted replicable genetic display package (rgdp) which displays said sbp member at the surface of the package, nucleic acid encoding said sbp member or a polypeptide component thereof being contained within the host cell in a form that is capable of being packaged using said rsdp component, whereby the genetic material of the rgdp displaying said sbp member encodes said sbp member or a polypeptide component thereof; said host organism being a mutator strain which introduces genetic diversity into the sbp member to produce said mixed population.

13. A method of producing a member of a specific binding pair (sbp), which method comprises:

expressing in recombinant host cells nucleic acid encoding said sbp member or a genetically diverse population of that type of sbp member, wherein the or each said sbp member or a polypeptide component thereof is expressed as a fusion with a component of a secreted replicable genetic display package (rgdp) which displays said sbp member in functional form at the surface of the package, nucleic acid encoding said sbp member or a polypeptide component thereof being

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contained within the host cell in a form that is capable of being packaged using said rgdp component, whereby the cenetic material of the rgdp displaying an sbp member encodes said sbp member or a polypeptide component thereof, said fusions being with bacteriophage capsid protein and the rgdps being formed with said fusions in the absence of said capsid protein expressed in wild-type form.

A method of producing a member of a specific binding pair (sbp) which method comprises: 10 expressing in recombinant host cells nucleic acid encoding said sbp member or a genetically diverse population of that type of sbp member, wherein the or each said sbp member or a polypeptide component thereof is expressed as a fusion with a component of a secreted replicable genetic display 15 package (rgdp) which displays said sbp member at the surface of the package, nucleic acid encoding said sbp member or a polypeptide component thereof being contained within the host cell in a form that is capable of being packaged using said rgdp component, whereby the genetic material of the rgdp displaying an sbp member encodes said sbp member or a polypeptide component thereof, said sbp member or polypeptide component thereof being expressed

- from a phagemid as a capsid fusion, and a helper phage, or a plasmid expressing complementing phage genes, is used 25 along with said capsid fusions to package the phagemid nucleic acid.
- A method according to claim 14, wherein said capsid protein is absent, defective or conditionally defective in 30 the helper phage.
- A method according to any one of claims 13 to 15 wherein the host cell is a mutator strain which introduces denetic diversity into the sbp member nucleic acid. 35
 - 17. A method according to any one of claims 9 to 16 wherein said library or genetically diverse population is obtained from:
 - the repertoirs of rearranged immunoglobulin genes (i) of an animal immunised with complementary sbp member.
 - the repertoire of rearranged immunoglobulin genes (ii)of an animal not immunised with complementary sbp member.
 - a repertoire of an artificially rearranged (iii) immunoglobulin gene or genes,
 - a repertoire of an immunoglobulin homolog gene (iv) or genes, or
 - a mixture of any of (i), (ii), (iii) and (iv). (v)
 - 18. A method according to any one of the preceding claims wherein said sbp member comprises a domain which is, or is homologous to, an immunoglobulin domain.
- 55 A method according to any one of the preceding claims

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wherein the rgdp is a bacteriophage, the host is a bacterium, and said component of the rgdp is a capsid protein for the bacteriophage.

- 20. A method according to claim 19 wherein the phage is a filamentous phage.
 - 21. A method according to claim 20 wherein the phage is selected from the class I phages fd, M13, fi, IfI, lke, $\rm ZJ/Z$. Ff and the class II phages Xf, Ffl and $\rm Pf3$.
 - 22. A method according to claim 20 or claim 21 wherein said sbp member or polypeptide chain thereof is expressed as a fusion with the gene III capsid protein of phage fd or its counterpart in another filamentous phage.
- 23. A method according to claim 22 wherein said sbp member or polypeptide chain thereof is inserted in the N-terminal region of the mature capsid protein downstream of a secretory leader peptide.
 - 24. A method according to any one of claims 19 to 23 wherein the host is E.coli.
- 25 25. A method according to any one of the praceding claims wherein nucleic acid encoding an sbp member polypeptide is linked downstream to a viral capsid protein through a suppressible translational stop codon.
- 30 26. A method according to any one of the praceding claims wherein the rgdps formed by said expression are selected or screened to provide an individual sbp member or a mixed population of said sbp members associated in their respective rgdps with nucleic acid encoding said sbp member or a polypeptide chain thereof.
 - $27.\,$ A method according to claim 26 wherein the rgdps are selected by affinity with a member complementary to said sbp member.
 - 28. A method according to claim 27 which comprises recovering any righs bound to said second member by washing with an eluant.
- 45 29. A method according to claim 28 wherein the eluant contains a molecule which competes with said rgdp for binding to the complementary sbp member.
 - 30. A method according to any one of the claims 27 to 29 wherein the rgdp is applied to said complementary sbp member in the presence of a molecule which competes with said package for binding to said complementary sbp member.
- 31. A method according to any one of claims 26 to 30, wherein nucleic acid derived from a selected or screened rydp is used to express said sbp member or a fragment or

derivative thereof in a recombinant host organism.

32. A method according to claim 31 wherein nucleic acid from one or more rgdps is taken and used to provide encoding nucleic acid in a further said method to obtain an individual sbp member or a mixed population of sbp members, or encoding nucleic acid therefor.

- 33. A method according to claim 31 or claim 32 wherein the expression end product is modified to produce a derivative thereof.
- 34. A method according to any one of claims 31, 32 and 33 wherein the expression end product or derivative thereof is used to prepare a therapeutic or prophylactic medicament or a diagnostic product.
- 35. Recombinant host cells harbouring a library of nucleic acid fragments comprising fragments encoding a genetically diverse population of a type of member of a specific binding pair (sbp), each sbp member or a polypeptide component thereof being expressed as a fusion with a component of a secretable replicable genetic display package (rgdp), so that said sbp members are displayed on surface of the rgdps in functional form and the genetic material of the rgdps encode the associated sbp member or a polypeptide component thereof.
- 36. Recombinant host cells according to claim 35, wherein said type of sbp member are immunoglobulins or immunoglobulin homologs, a first polypeptide chain of which is expressed as a said fusion with a component of the rgdp and a second polypeptide chain of which is expressed in free form and associates with the fused first polypeptide chain in the rgdp.
 - 37. A helper phage whose genome lacks nucleic acid encoding one of its capsid proteins, or whose encoding nucleic acid therefor is conditionally defective, or which encodes said capsid protein in defective or conditionally defective form.

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- 38. A bacterial host cell containing a filamentous phage genome defective for a capsid protein thereof and wherein the host cell is capable of expressing capsid protein complementing said defect such that infectious phage particles can be obtained therefrom.
- 39. A bacterial host cell according to claim 38 wherein said complementing capsia protein is expressed in said host from another vector contained therein.
- 40. A bacterial host cell according to claim 38 or claim 39 wherein the defective capsid protein is gene III of phage fd or its counterpart in another filamenrous phage.

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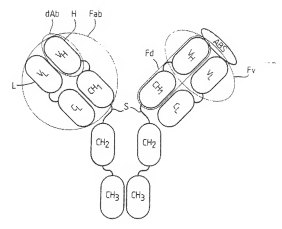
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- 41. Recombinant E.coli TG1 M13K07 gIII No. 3 (NCTC 12478).
- 42. A phage having the form of a replicable genetic display package displaying on its surface in functional form a member of a specific binding pair or a binding domain thereof.
 - 43. A kit for use in carrying out a method according to any one of claims 1 to 34, said kit including:
 - (i) at least one vector having an origin of replication for single-stranded bacteriophage, a restriction site for insertion of nucleic acid encoding said sbp member or a polypeptide component thereof in the 5' end region of the mature coding sequence of a phage capsid protein, and with a secretory leader sequence upstream of said site which directs a fusion of the capsid protein and sbp polypeptide to the periplasmic space of a bacterial host; and
 - (ii) ancillary components required for carrying out the method.

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Fig. 1.



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Fig.2(i)

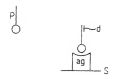
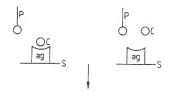
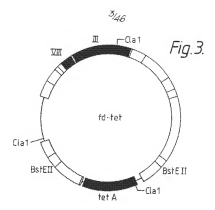


Fig.2(ii)





WO 92/01047 PCT/GB91/01134



fd-tet

cleave with BstEII

fill in with Klenow

re-ligate

f FDT6Bst

in vitro mutagenesis (oligo 1)

fFDTPs/Bs

in vitro mutagenesis (oligo 2)

FDTPs/Xh
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Fig. 4.1

ACA ACT TTC AAC AGT TGA GGA GAC GGT GAC CGT AAG CTT CTG CAG TTG GAC CTG AGC GGA GTG AGA ATA (1620) Cffgo 1 Oligo 2

ACA ACT TTC AAC AGT TTC CCG TTT GAT CTC GAG CTC CTG CAG TTG GAC CTG

Oligo 3

GTC GTC TTT CCA GAC GTT AGT

GENE III

CLEAVAGE SITE SIGNAL

FIG. 4.2

A TCT CAC TCC GCT

8 TCT CAC TCC GCT CAG GTC CAA <u>CTG CAG</u> AAG CTT AC<u>G GTC ACC</u> GTC TCC TCA ACT GTT GAA AGT Pst I 0 / 0 / 0

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GAA ACT GTT GAA AGT C TOT CAC TOO GOT CAB GTO CAA CITG CAB GAG CITC GAB ATO AAA CGB

GAA ACTGTT GAA AGT

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GENE III

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Fig. 5.

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Fig.5 cont.

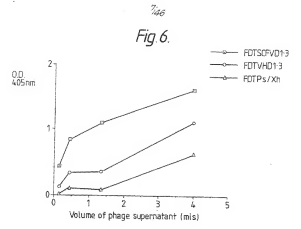
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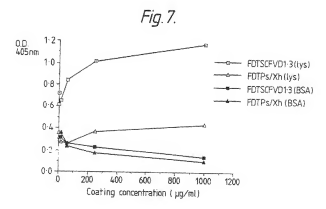
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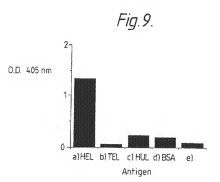


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Fig. 8.





9₄₆ Fig.10.

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G P G L V A P S Q S L S I T C T V S G F GGACCIGGCCTGGTGGCCCCTGCCACGCCTCCACGGTTC L30 140 150 160 170 180

L G M I W G D G N T D Y N S A L K S R L CTGGGAATGATTTGGGGTGATGAAACACAGAGTATTAATTCAGGTGCTAAATCCAGACTG 250 260 270 280 290 300

S I S K D N S K S Q V F L K M N S L H T AGCATCAGCAACGCCAACGCCCAACGTTTTCTTAAAAATGAACAGTCTGCACACT 310 320 330 340 350 360

Q G T T V T V S S A S T K G P S V F P L CAAGGCACCACGCTCACGTCCACGCACGACGACGCACGACGTCCTCCCCCC 430 440 450 460 470 480

A P S S K S T S G G T A A L G C L V K D GCACCCTCCTAAAAACCACCTCTGGGGCACACGGGCCCTGGGTCAAGACAC 490 500 510 520 530 540

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Fig.10 cont.(1)

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850 860 870 880 890 900 L S A S V G E T V T I T C R A S G N I H CCCTTCTCGCTCCGTCGCAAACTCTCACATCACATCTCCAAATATTC 910 920 930 940 950 960 N Y L A W Y Q Q K Q G K S P Q L L V Y Y																									
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Fig.10 cont.(2)

TTTLADGVPSRFSGSGSGTO ATTACAACAACCUTTAGCAGATGGTGTGCCATCAACGTTTCAGTGGCAGTGGGATCAGGAACAC 1030 1040 1050 1060 1070 1080

YSLKINSLOPEDFGSYYCOH DAKTITIDAHIRATEKATATITITIAKAAHITYSAYSITYYSAANIMADAATITITATIKA 1090 1100 1110 1120 1130 1140

FWSTPRTFGGGTKLEIKRTV ATTITIGACIACIOCIOGACGITIGGICGACCCACCOCACCIVUACATCAACCGAC 1150 1160 1170 1180 1190 1200

AAPSVFIFPPSDEQLKSGTA TOGCTGCACCATCTGTCTTCATCTTCCCCCCATCTGATGACCACTTGAAATCTGGAACTG 1210 1220 1230 1240 1250 1260

SVVCLLNNFYPREAKVOWKV CCITCIGITGIGIGCCIICCIGAADAACHTCDATCCCAGAGAGGCCCAAACHACAGTICAACG 1270 1280 1290 1300 1310 1320

D N A L Q S G N S Q E S V T E Q D S K D TOCATIAACGCOUTOCAATOGGGTAACTOCCACGAGAGTGTCACAGAGCAGCAGCAGCAGCAGC 1330 1340 1350 1360 1370 1380

STYSLSSTLTLSKADVRKHK ACAGCACCTIACAGCCTCAGCAGCACCACCCTGACCCAAACCAGACTACGAGAACACA 1390 1400 1410 1420 1430 1440

V Y A C E V T H Q G L S S P V T K S F N AAGICTAGGCCTGGGAAGTCACCCATCAGGCCCTTGAGGTGGCCCGTCACAAAGAGCTTTCA 1460 1470 1480 1450 1490 3500

R G E S * * ACCIGCIGAGACITCATACITAACAATTIC 1510 1520



Fig.10 cont.(3)



FabD1.3 in pUC19

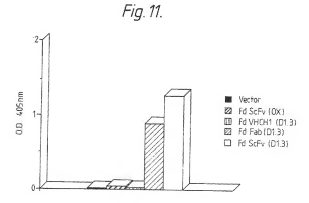




Fig.12.



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Fig.13.

L O G G L V Q P G G G CAG GTG CAG CTG CAG GAG TCA GGA GGA GGC TTG GTA CAG CCT GGG GGT PstI TOT CTG AGA CTC TCC TGT GCA ACT TCT GGG TTC ACC TTC AGT AAT TAC Y M G W V R Q P P G K A L E W L TAC ATG GGC TGG GTC CGC CAG CCT CCA GGA AAG GCA CTT GAG TGG TTG N G Y T GGT TCT GTT AGA AAC AAA GTT AAT GGT TAC ACA ACA GAG TAC AGT GCA S V K G R F T I S R D N TOT GTG AAG GGG CGG TTC ACC ATC TCC AGA GAT AAT TTC CAA AGC ATC INTLRTED SA CTC TAT CTT CAA ATA AAC ACC CTG AGA ACT GAG GAC AGT GCC ACT TAT Y Y TAC TOT GCA AGA GGC TAT GAT TAC GGG GCC TGG TTT GCT TAC TGG GGC L V T V S S g g g s g g g s T CAA GGG ACC CTG GTC ACC gtc tcc tca ggtggoggcggttcaggcggaggtggctct L T 9 g g g * d í B 0 L 99-99199-99010998C atc GAG CTC ACC CAA ACT CCA CTC TCC CTG CCT GTC SacI S 8 AGT CTT GGA GAT CAA GCC TCC ATC TCT TGC AGA TCT AGT CAG AGC ATT V H S N G N T Y L EWYLQKP GTA CAT AGT AAT GGA AAC ACC TAT TTA GAA TGG TAC CTG CAG AAA CCA SPKLL I Y K V S GGC CAG TOT COA AAG CTC CTG ATC TAC AAA GTT TCC AAC CGA TIT TOT GGG GTC CCA GAC AGG TTC AGT GGC AGT GGA TCG GGG ACA GAT TTC ACA LKISRVEAEDLGV Y Y C CTC AAG ATC AGC AGA GTG GAG GCT GAG GAT CTG GGA GTT TAT TAC TGC SHVPYTPGGGTKL TIT CAA GGT TCA CAT GIT CCG TAC ACG TIC GGA GGG GGG ACC AAG CTC E I K GAG ATC AAA CGG XhoT

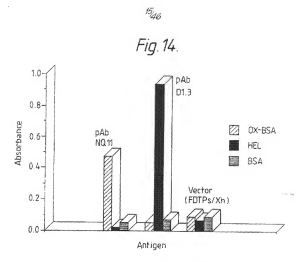


Fig. 15.

5¹ END R T P E M P V L
TCT CAC A<u>GT GCA C</u>AA ACT GTT GAA CGG ACA CCA GAA ATG CCT GTT CTG
Apal?

S¹END

K. A. A. L. G. L. K.

AAA GCC GCT CTG GGG CTG AAA <u>GCG GCC GC</u>A GAA ACT GTT GAA AGT etc.

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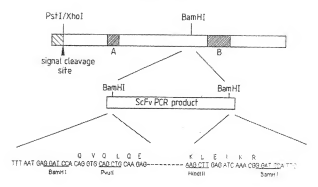
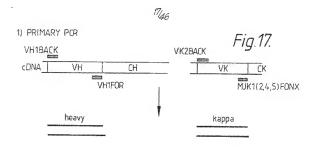
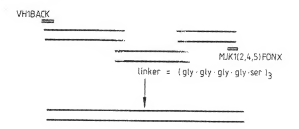


Fig.16(2)

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2) ASSEMBLY PCR



3) ADDING RESTRICTION SITES
VHBKAPA10

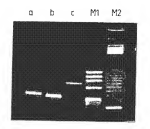
JK1(2,4,5)NOT10

Apa L1

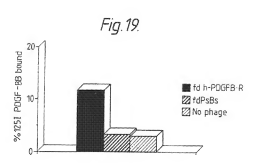
Not1

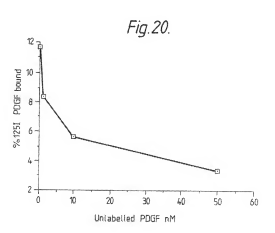


Fig.18.



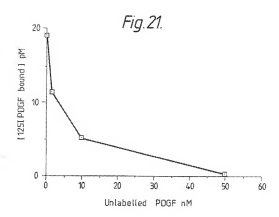


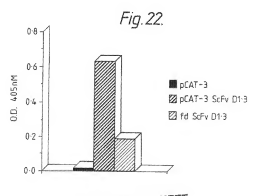




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Fig.23.

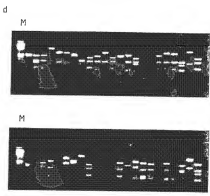


Fig. 24.

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~ % # O O M M O D P	*	*************

VH sequences

Fig. 24 cont.

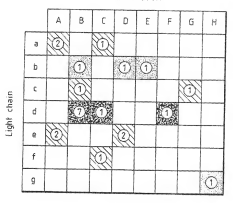
Vx sequences

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0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	9x cox 1 cox 1 1 ke 2 cox 1 ke 2
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CD#1 GABGBT807L8 SABSBYBBTLN BAGBBT808TLB BAGBST808TLB BAGBSTBTB GASSBTRTP06 GASSBTRTP06	8A SEGURE S. RA CASE C. RA
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Fig. 25.

HEAVY CHAIN



00405nm in ELISA









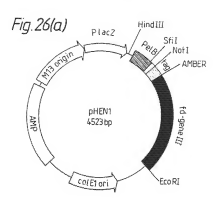


Fig.26(b)

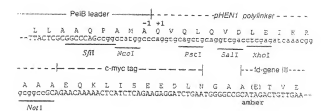
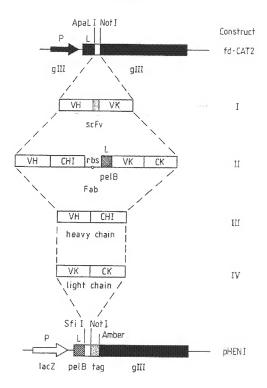




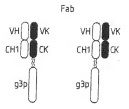
Fig. 27.

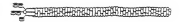


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Fig. 28.

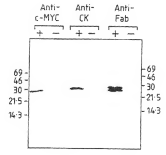


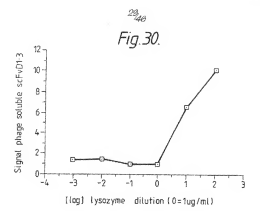


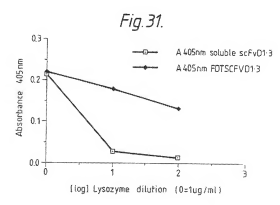


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Fig.29.







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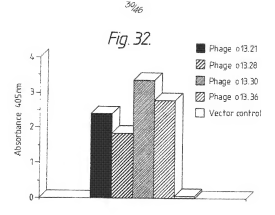
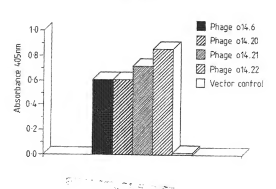


Fig. 33.

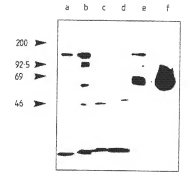


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Fig. 34.



³²46 Fig. **3**5.

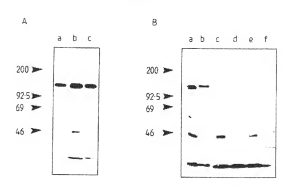
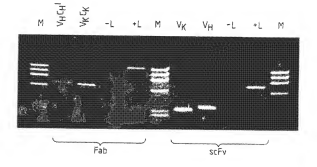
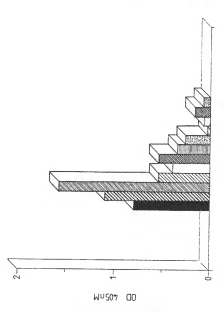


Fig. 36.



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Fig. 37.

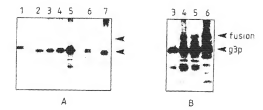


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Fig.38.



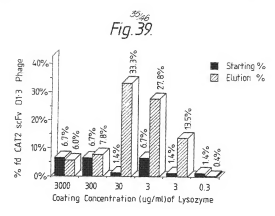
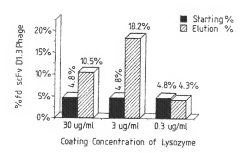


Fig. 40.



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Fig. 41.

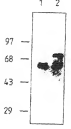
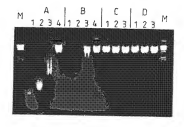


Fig. 42.



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Fig. 43.

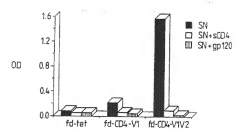


Fig. 44 (1)

TICTATTCTCACAGIGGENOAGGIGCAGCIGCAGCAGTGGGGGCIGAGCTIGIGAAGCCIGGGGCTICAGIGAAGCIGICCIGCAAGGCI angatangagicicocgiisiocaggicgacgicgicagacccggacicgaacticgaacccgaagicgaacciicgaacgiiccga PheTyrSerHisSerhla3inVaiGinLeuGinGinSerGlyAlaGiuLeuVaiLysProGlyAlaSerValLysLeuSerCysLysAla

tetggctacaccticaccaggtactggatgcactgggtgaagcagaggcttggacgaggceitgagtggattggattggattgatgcttat AGACCGATGTGGAAGTGGTGGATGACCTACGTGACCACTTCGTCTCGGACCTGCTGGTGCTGCACCTAACCTTCCTAACTAGGATA SerglyTyrThrpheThrSerTyrTrpMetHisTrpValLysGlnArgProSlyArgGlyLeuGluTrp11eGlyArgIleAspProAsn 140 130 120 330

agtiggtiggtactaagtacaatgagaagticaagaggaaaggcacactgagctstagacacatcctccagcacagcctacatgcac SerglyglythriystyrasaglulysPhelysSerlysAlathrleuthrvalasplysProserserthralyrMetGlnLeuser 210

AGCCTGACATCTGAGGACTCTGCGGTCTNTTATTGT@AAGNIACGACTACGTAGTAGCTACTACTTTGACTACTGGGGGCCAAGGGAAC TCEGACTETAGACTECTGAGACGCCAGATAATAACACGTTCTATGCTGATGCCATCATCGATGATGAAAGAGATGACCCGGGTTCCCTGG SerlauthrSergluaspSerAlaValTyrTyrCys<u>Ala</u>ArgTyrAspTyrGlySerSerTyrTyrPheAspTyrTrpGlyGlnGlyThr 340

acgstcaccetctcctcmsgtgsaggcggttcaggsgaggtggctctggcggtggcgsatcccaggctgttcggatctaggatctgca TGCCAGTGGCAGAGGAGTCCACCTCCGCCAAGTCCGCCTCCACGGAGACCGCCAACGCCTAGGGTCCGACAACCCTGTGTCCTTAGACGT ThrvalThrvalserserblyglyglyglyserglyglyglyglyserglyglyglyglyglyglyglysexblnalavalglyThrglnglusernia 430 410 380

CTCACCACATCACCTGGTGAAACAGTCACACACATGTCGCTCAAGTACTSGGGGTGTTACAACTAGTAACTATGCCAACTGGGTCGAA gagtiggtetagtiggaccaltttetcagtigagtgaacagcgagttlatgaccccgacaatgatgatgatgatacogttgacccaaggt Leuthrthrserfroglygluthrvalthrleuthrcyzdrgsersefthrslydlavalthrthrserdantyrdladantrpvalgln

gharaccasatcattattattcactggtctaataggtggtaccaacaaccaggtctccagststtcctgccagattttcaggctccctgatt CTTTTEGETCINGTANATAAGTGACCAGATTATCCACCATGGTTGTTGGCTCGAGGTCCALAAGGACGGTCTAAGAGTCCGAGGGACTAA GlulysPrcAspHisLeuPheThrGlyLeuIleGlyGlyThrAsnAsnArgAlaFrcGlyValPrcAlaArgPheSerflySerLeuIle

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Fig. 44 (ii)

GlyAsplysAlaAlaLeuThrIleThrGlyAlaGlnThrGluAspGluAlaIleTyrPheCysAlaLeuTrpTynberAsnHisTrpVal



Fig.45.

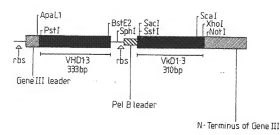
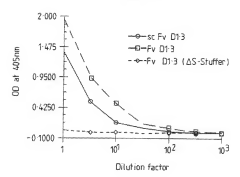


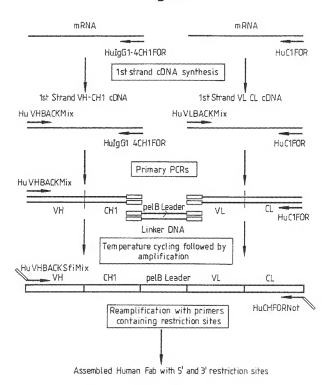
Fig. 46.



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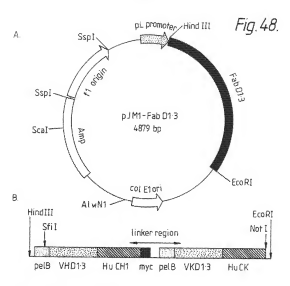


Fig. 47.



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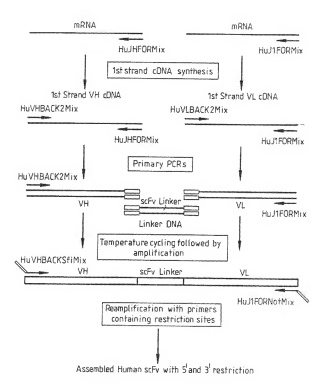




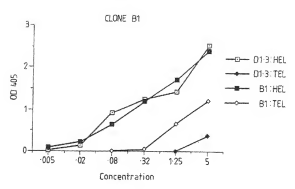
C. Sequence of linker region

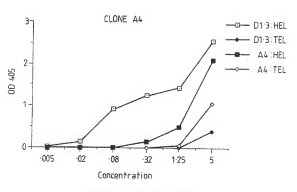


⁴³/₄₆ Fig.49.









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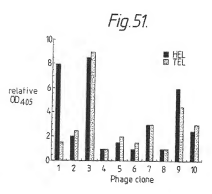


Fig.53.

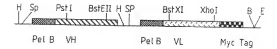


Fig. 52.

DIELTQSPALMAASPGEKVTITCSV888I888NLHWYQQKSETSPKPWIYGTSNLAS D1.3 DIQMTQSPASLSASVGETVTITCRASGNIHNYLA WYQQKQGKSPQLLVYYTTILAD DIELTQSPSSLSASLGERVSLTCRASQDIGSSLN WLQQEPDGTIKRLIYATSSLDS CDR 2 MIF M2.1

CDR 3

GVPSRFSGSGSGTQYSLKINSLQPEDFGSYYCQHFWSTPRTFGGGTKLEIKR GVPKRFSGSRSGSDYSLTISSLESEDFVDYYCLQYASSPWTFGGGTKLELKR MIR

GVPVRFSGSGSGTSYSLTISSMEAEDAATYYCQQWSSYPLTFGAGTKLEIKR

M2.1

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INTERNATIONAL SEARCH REPORT

PCT/GB 91/01134

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See the whole document, especially Figu	See the whole document, especially Figure 1			
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